



Understanding Small Separators in Road Networks

Master's Thesis of

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I declare that I have developed and written the enclosed thesis completely by myself. I have not used any other than the aids that I have mentioned. I have marked all parts of the thesis that I have included from referenced literature, either in their original wording or paraphrasing their contents. I have followed the by-laws to implement scientific integrity at KIT.

Karlsruhe, July 1, 2025

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(Samuel Born)

Abstract

Zusammenfassung

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1 Introduction

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1.1 Motivation

In graph theory, a separator is a subset of vertices whose removal divides the graph into disconnected components of roughly equal size. The size of these separators significantly affects the performance of numerous algorithms, particularly those utilizing divide-and-conquer approaches. For instance, Tarjan's and Lipton's foundational work on planar graphs [LT77] demonstrates their utility in optimizing algorithms for many problems like e.g. maximum flow.

Empirical studies suggest road networks have balanced separators on the order of $\mathcal{O}(n^{1/3})$ [DSW16], which is significantly smaller than the $\mathcal{O}(n^{1/2})$ worst-case bound for planar graphs [LT79]. This is noteworthy given that road networks are can be treated as nearly planar due to their geographic nature (few crossings from overpasses/tunnels).

In road networks, these separators enable the creation of effective node orders, which are critical for the performance of search queries in advanced routing algorithms like CCH. This thesis seeks to uncover the properties responsible for the presence of such small separators in road networks. We aim to determine whether these separators stem from inherent graph characteristics, such as limited vertex degrees or sparsity, or from physical real-world features, such as borders, rivers, or a hierarchical structure. Gaining insight into these properties promises to advance our theoretical understanding and offers practical benefits, such as identifying new applications or generating comparable synthetic graphs.

Road networks represent an intriguing subject for the study of graph separators. Classical results within graph theory primarily establish balanced separators characterized by asymptotic sizes such as $\Theta(1)$, common in structures like trees, or $\Omega(\sqrt{n})$, e.g. planar graphs and unit disk graphs. To the best of our knowledge, established graph classes consistently exhibiting separator sizes strictly between these $\Theta(1)$ and $\Omega(\sqrt{n})$ bounds are not prominently featured in the graph theory literature. This finding positions road networks within this sparsely populated intermediate complexity range, thereby highlighting the compelling nature of investigating their structural properties.

1.2 Contribution

todo

1.3 Outline

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2 Preliminaries

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2.1 Graph Theory

Road networks can be modeled as graphs. A graph G is formally defined as a pair (V, E) , where V represents a finite set of vertices (or nodes) and E represents a set of edges connecting pairs of vertices. In many applications, particularly route planning, graphs are augmented with a weight function $w : E \rightarrow \mathbb{R}^+$, assigning a positive real value such as distance or travel time to each edge. However, for the purpose of this thesis, the topological structure of the graph is of primary interest, and we will not focus on edge weights. We will also only consider simple graphs, meaning graphs without multiple edges between the same pair of vertices and without edges connecting a vertex to itself (loops). Furthermore, as the concept of separators primarily applies to connectivity, we will consider undirected graphs, where edges represent symmetric relationships. An edge connecting vertices u and v in an undirected graph is denoted as the set $\{u, v\}$. The neighborhood of a vertex v is defined as the set of vertices adjacent to v , denoted as $N : V \rightarrow \mathcal{P}(V)$.

A graph *embedding* assigns each vertex $v \in V$ of a graph $G = (V, E)$ to a unique point p in a specific geometric space, such as the Euclidean plane \mathbb{R}^2 or the surface of a sphere. We then consider edges as straight line segments connecting the points corresponding to their incident vertices.

Throughout this thesis, the term *graph size* refers specifically to the number of vertices, $|V|$. We frequently adopt the notation n for the number of vertices, $|V|$, and m for the number of edges, $|E|$. It is worth noting that for many graph classes discussed herein, particularly sparse graphs like road networks, the number of edges m is asymptotically linear in the number of vertices n . For planar graphs, a specific bound guarantees this linear growth. Euler's formula for connected planar graphs states that $n - m + f = 2$, where f is the number of faces (regions) defined by the graph embedding. By observing that each face is bounded by at least three edges (for $n \geq 3$) and each edge separates at most two faces, we derive the inequality $2m \geq 3f$. Substituting $f = 2 - n + m$ from Euler's formula into this inequality yields $m \leq 3n - 6$. This confirms the linear relationship between the number of edges and vertices for planar graphs. Therefore, the choice of n as the measure of size generally does not impact asymptotic complexity results for the graphs under consideration.

2.2 Graph Separators

A vertex separator (or simply separator) of a graph $G = (V, E)$ is a subset of vertices $S \subseteq V$ whose removal disconnects the graph into two or more components. More formally, the subgraph induced by $V \setminus S$, denoted $G[V \setminus S]$, is disconnected. For algorithmic applications, particularly divide-and-conquer strategies, balanced separators are crucial.

Let V_1, \dots, V_k be the vertex sets corresponding to the connected components of the subgraph $G[V \setminus S]$. Most often, the removal of such a separator yields exactly two components ($k = 2$), as partitioning the graph into a larger number of components generally demands a larger separator. For a given constant $\alpha \in (0, 1)$, a separator S is termed α -balanced if the size of every resulting component V_i is bounded. Specifically, the condition $|V_i| \leq \alpha|V|$ must hold for all $i \in \{1, \dots, k\}$. A simple illustration of a balanced separator is shown in Figure 2.1. A common requirement is $2/3$ -balancedness, meaning each component contains at most $2/3$ of the original graph's vertices. Balancedness ensures that recursive applications of the separator lead to subproblems of substantially smaller size, which is essential for the efficiency of algorithms based on this technique.

Furthermore, minimizing the size of the separator S itself is critical for algorithmic performance. The size of the separator is typically evaluated asymptotically as a function of the number of vertices $n = |V|$ e.g. n^β for $\beta \in (0, 1)$.

The concept of *recursive* α -balanced separators extends this idea by ensuring that the property of finding small, balanced separators persists in the resulting subgraphs. Specifically, after removing an α -balanced separator S from G , each induced subgraph $G[V_i]$ (for $i = 1, 2, \dots, k$) can itself be partitioned using another α -balanced separator of small size.

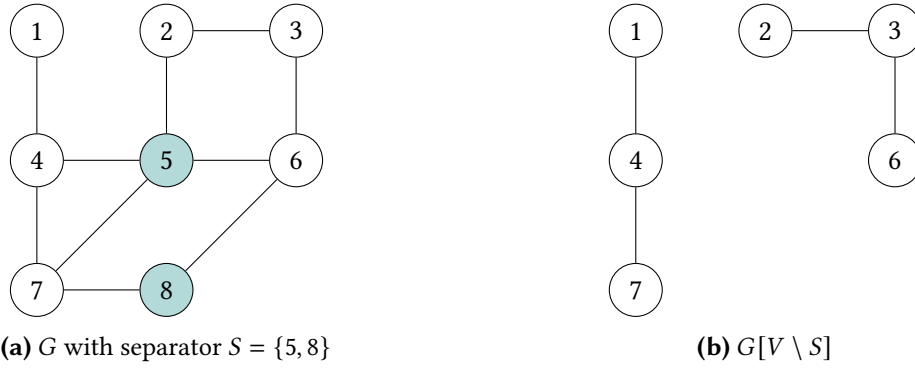


Figure 2.1: Example of a well balanced separator in a graph. The vertices 5 and 8 form a balanced separator that disconnects the graph into two components.

To compute separators, various algorithms can be employed. In this thesis, we primarily utilize InertialFlowCutter [GHUW19]. This algorithm leverages geometric embeddings, often available for road networks, to compute high-quality node orderings efficiently. These node orderings serve as the basis for extracting separators from the graph using the method described below [Blä+25]. Employing this combined approach provides a practical pathway to generate separators for our analysis, adapting the use of InertialFlowCutter's output to suit the specific requirements of this work. However, InertialFlowCutter requires such a geometric embedding as input. For graphs without an embedding, we utilize the KaHIP (Karlsruhe High Quality Partitioning) framework [SS13].

When using InertialFlowCutter, the resulting node ordering is interpreted as an elimination order for the vertices of the graph $G = (V, E)$. Based on this order, a chordal supergraph $G_C = (V, E \cup F)$ is constructed, where F represents the fill-in edges. The chordal supergraph is constructed by processing vertices v according to their rank. Fill-in edges are added such that for each vertex v , all its neighbors with a rank greater than $rank(v)$ form a clique. An efficient implementation connects the neighbor u_{min} with the minimum rank among those

where $\text{rank}(u_{\min}) > \text{rank}(v)$ to all other neighbors w also satisfying $\text{rank}(w) > \text{rank}(v)$. This suffices because the responsibility for adding edges between the remaining pairs of these higher-ranked neighbors w is effectively delegated to u_{\min} .

Afterwards, a tree structure T can be constructed. Each node $v \in V$ selects its parent in the tree as the neighbor u that appears earliest in the elimination order among all neighbors w in G_C with $\text{rank}(w) > \text{rank}(v)$. If a node has no neighbors later in the order, it becomes the root.

Separators in the original graph G can be derived from this tree structure using a traversal algorithm. The fundamental idea is to identify paths representing non-branching segments of the tree. Starting from a node r (representing the current subgraph), the traversal follows a path $P = (v_1 = r, v_2, \dots, v_k)$ downwards, where each node v_i ($1 \leq i < k$) has exactly one child v_{i+1} in the tree. The path ends at node v_k , which is the first node encountered that does not have exactly one child (i.e., it has zero or multiple children). The set of vertices on this path, $S = \{v_1, v_2, \dots, v_k\}$, forms a separator. Its size is k , the number of nodes on the path. The traversal algorithm continues recursively into these subtrees. An overview of this process is illustrated in Figure 2.2.

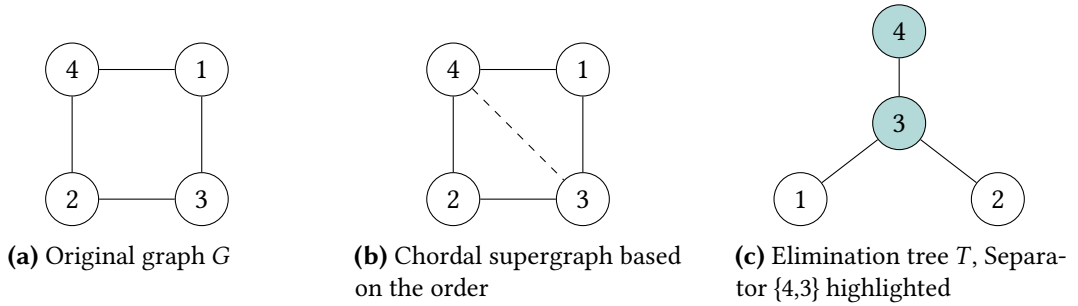


Figure 2.2: Example Process of deriving a separator from a node order. Node labels indicate their rank in the node order.

To ensure separators yield balanced partitions, the extraction process is refined using a 'significance threshold' based on relative subgraph size. Child nodes are only considered 'significant' if the subgraph they represent exceeds this threshold (e.g., contains at least 5% of the nodes in the parent's subgraph). When tracing a potential separator path $P = (v_1, \dots, v_k)$ downwards, the path extends from v_i to v_{i+1} only if v_{i+1} is the *single* significant child of v_i . The path $S = \{v_1, \dots, v_k\}$ is finalized as the separator upon reaching the first node v_k that possesses *two or more* significant children. This ensures separators correspond to meaningful branches in the tree structure concerning substantial graph parts. Figure 2.3 illustrates an example of this process.

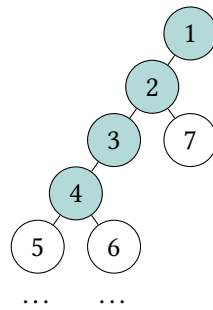


Figure 2.3: Separator identification with a significance threshold. The path extends downwards from node 1. At node 2, child 7 represents an insignificant subgraph (below the threshold), so the traversal continues via the single significant child path towards node 3. Node 4 is the first node encountered with two children (5 and 6) that both represent significant subgraphs. Therefore, the process stops here, and the identified separator is $S = \{1, 2, 3, 4\}$.

3 CCH: A State-of-the-Art Application

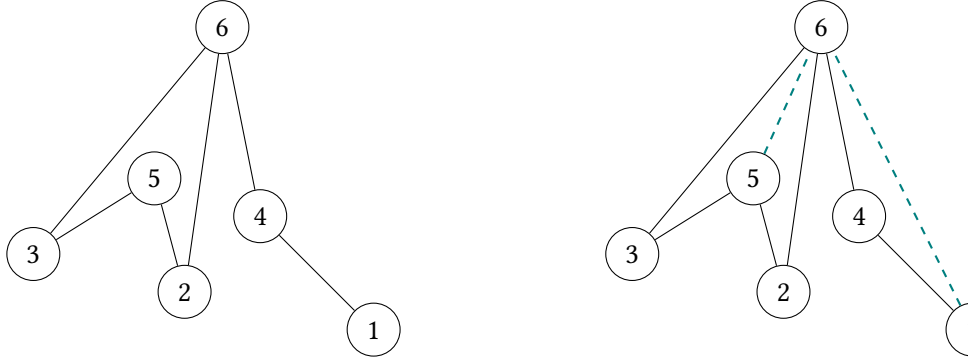
As established in the introduction, real-world road networks exhibit small separators. This observation, stemming from experimental results of the Customizable Contraction Hierarchies (CCH) paper [DSW16], serves as a key inspiration for this thesis. This chapter delves into CCH as a state-of-the-art route planning technique to illustrate precisely how such small separators are leveraged to achieve high query performance, particularly in dynamic scenarios where edge weights, such as those reflecting traffic conditions, frequently change. Understanding this application provides further context for the significance of the small separator phenomenon that this thesis investigates.

Customizable Contraction Hierarchies address the challenge of dynamic edge weights by employing a three-phase approach: an initial, topology-dependent precomputation; a subsequent, fast customization phase that incorporates current edge weights; and finally, an efficient query phase [DGPW11 | DSW16]. The core idea underpinning CCH involves strategically inserting shortcut edges into the graph, analogous to the concept used in the original Contraction Hierarchies (CH) algorithm [GSSD08]. These shortcuts bypass sequences of original edges, effectively contracting the graph and speeding up queries. The following provides an overview of the CCH algorithm, with a focus on how its components benefit from the small separators found in road networks.

Precomputation The CCH precomputation phase introduces shortcut edges based on a given vertex order. These shortcuts effectively bypass sections of the graph, allowing algorithms to skip over entire subgraphs, unless the target node resides within such a subgraph. Furthermore, the specific process of inserting shortcuts based on the contraction order guarantees that any shortest path in the original graph corresponds to an 'up-down' path in the hierarchy defined by the vertex ranks [GSSD08]. An 'up-down' path consists of a sequence of edges leading to vertices with increasing ranks (the 'up' segment), followed by a sequence of edges leading to vertices with decreasing ranks (the 'down' segment). This property enables efficient bidirectional search by restricting exploration to higher-ranked neighbors.

This order is defined by a bijection $\pi : \{1, \dots, n\} \rightarrow V$, where $n = |V|$. We will call the inverse of this order rank : $V \rightarrow \{1, \dots, n\}$, which assigns each vertex its position in the order. The core process involves iteratively contracting vertices in order of their rank, starting from rank 1 up to n . Contracting a vertex v_i involves removing it and its incident edges from the current graph representation. For every pair of (higher ranked) neighbors $u, w \in N(v_i)$, a shortcut edge (u, w) is introduced. Resulting multi-edges are simplified. We call the resulting graph $G_C = (V, E_C)$, where $E_C = E \cup F$ and F represents the set of shortcut edges. The contraction process is illustrated in Figure 3.1.

A primary objective when selecting the vertex order is to minimize the number of shortcut edges introduced during the contraction process. Minimizing shortcuts is beneficial for both storage and query efficiency [DSW16]. However, solely minimizing the number of added shortcuts may not be sufficient in all cases. Different heuristics for selecting the contraction order exist.



(a) Input graph. Already converted to be undirected and simple.

(b) Graph after precomputation, new shortcut edges are shown in teal.

Figure 3.1: Example of the CCH precomputation step. Nodes are named and positioned based on their rank.

Nested Dissection One method for computing good vertex orders are Nested Dissections. The process begins by identifying a small, balanced separator in the graph. Nodes within this separator are conceptually removed, partitioning the graph into smaller components. These separator nodes are designated as high-rank nodes in the hierarchy and are consequently placed towards the end of the final node ordering. This procedure is then applied recursively to the remaining components. Figure 3.2 provides a visual representation of this recursive partitioning strategy.

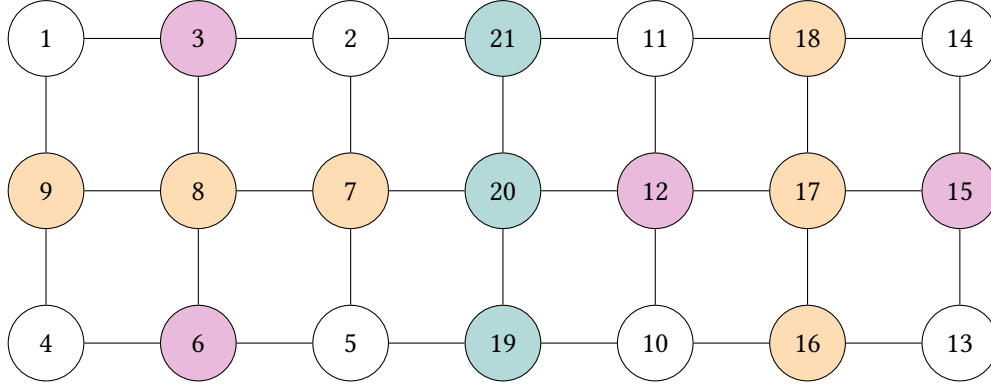


Figure 3.2: Example of a Nested Dissection. The top level separator is shown in teal, the second level in orange and the third level in purple. The nodes are named according to their rank in the resulting order.

Customization Customization assigns the current metric's weights to the original edges within the CCH supergraph G_C and initializes shortcut edge weights to infinity. Following this initialization, edge weights are systematically updated to ensure the triangle inequality holds throughout G_C .

To achieve this, the concept of a lower triangle is employed. Given an edge $\{x, y\} \in E_C$, a lower triangle is formed by the vertices $\{x, y, z\}$ if the edges $\{z, x\}$ and $\{z, y\}$ also exist, and $rank(z) < \min\{rank(x), rank(y)\}$. The customization algorithm iterates through the

vertices of the graph in ascending order of their precomputed rank. For each vertex x , it considers all upward edges $\{x, y\}$ in the graph, where y is a neighbor of x and $rank(y) > rank(x)$. For every such edge $\{x, y\}$ we determine all lower triangles $\{x, y, z\}$. If the path through z offers a shorter connection, the weight of the edge $\{x, y\}$ is updated to this smaller value: $w(x, y) \leftarrow \min\{w(x, y), w(x, z) + w(z, y)\}$. The detailed procedure is outlined in the pseudocode presented in Algorithm 3.1. An illustration of the customization process is provided in Figure 3.1.

Note that the outlined algorithm only considers undirected edge weights, the algorithm can be extended to directed edge weights. Details can be found in [DSW16].

Algorithm 3.1: CCH Customization

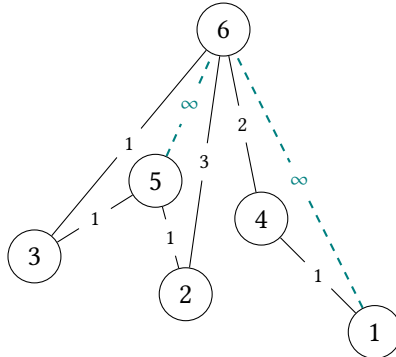
Input: $G_C = (V, E_C)$, node ordering π , edge weights w

Output: Customized CCH graph

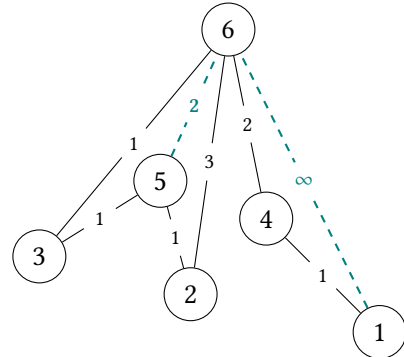
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1 forall  $x$  in  $V$  in ascending order of rank do
2   forall upward edges  $\{x, y\}$  in  $E_C$  do
3     forall lower triangles  $\{x, y, z\}$  associated with  $\{x, y\}$  do
4        $w(x, y) \leftarrow \min\{w(x, y), w(x, z) + w(z, y)\}$ 

```



(a) Graph after precomputation. Weights are added to the edges. Shortcuts get weight ∞ .



(b) Graph after customization. The shortcut edge $\{5, 6\}$ is updated to weight 2.

Figure 3.3: Example of the CCH customization step.

Query To answer a shortest path query between a source node s and a target node t , the algorithm utilizes a structure known as the elimination tree. The elimination tree is defined on the nodes of G_C . The parent of a node v in the elimination tree is the neighbor p of v in the CCH graph that has the lowest rank among all neighbors with a rank strictly greater than the rank of v . Figure 3.4 illustrates the elimination tree for the example graph shown in Figure 3.1. The query algorithm performs a bidirectional search upwards in this elimination tree, starting from s and t .

The core query process operates iteratively. Let u_s and u_t be the current nodes in the upward search originating from s and t , respectively; initially, $u_s = s$ and $u_t = t$. The algorithm proceeds until the root of the elimination tree is reached. In each step, the ranks of the current nodes u_s and u_t are compared. If u_s has a smaller rank than u_t , the algorithm relaxes all outgoing edges $\{u_s, v_i\}$ present in G_C . Subsequently, u_s is updated to become its

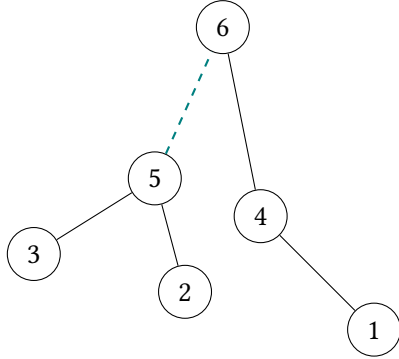


Figure 3.4: Elimination tree for the example graph in Figure 3.1.

parent node in the elimination tree. Otherwise (if u_t has a rank less than or equal to that of u_s), the algorithm relaxes all outgoing edges $\{u_t, v_i\}$ existing in the CCH graph. Following the relaxation step, u_t is updated to its parent in the elimination tree. This process continues, effectively exploring paths upwards towards higher-ranked nodes. The correctness of this query algorithm for computing shortest path distances has been established; a detailed proof, which is beyond the scope of this thesis, can be found in [DSW16].

Complexity The size of the separators found significantly impacts the efficiency of CCH queries. CCH queries restrict exploration to edges leading towards higher-ranked nodes (upward edges). Consider the separator identified at the highest level of the recursion, which contains approximately n^β nodes. When a query initiates within a component defined by this separator, nodes located in other components cannot be reached without traversing downwards through a separator node, violating the upward search constraint. This containment effect applies recursively within the sub-components generated during the nested dissection. Let α denote the balance factor. The sub-components at recursion level i consequently have size at most $\alpha^i \cdot n$. Analyzing the total bound of the search space involves summing these separator sizes across the finite levels i of the recursion. This sum can be bounded by approximating it with the corresponding infinite geometric series [BCRW16]:

$$\begin{aligned}
 & \sum_{i=0}^{\infty} (\alpha^i \cdot n)^\beta \\
 &= n^\beta \cdot \sum_{i=0}^{\infty} \alpha^{i \cdot \beta} \\
 &= n^\beta \cdot \frac{1}{1 - \alpha^\beta} \quad \text{Geometric series, since } \alpha \in (0, 1) \\
 &\in \mathcal{O}(n^\beta)
 \end{aligned}$$

This analysis demonstrates that the total search space size explored during a CCH query is bounded by $\mathcal{O}(n^\beta)$, under the assumption that small separators can be found recursively. Note that we do not have worst case guarantees for the size of the separators in real road networks, but we can expect them to be small in practice. Thus, the performance of the CCH algorithm is directly linked to the ability to find small separators.

4 Road Network Properties

This chapter provides an empirical analysis of several fundamental properties of real-world road networks. We investigate key characteristics such as separator sizes, degree distribution, planarity, hierarchy, and diameter to establish a baseline understanding of these networks. These empirical findings serve as a foundation for the synthetic graph generation and analysis presented in subsequent chapters.

4.1 Separator Sizes

To empirically investigate the relationship between graph size and separator size in road networks, we analyze the DIMACS Europe provided by PTV [PTV09]. We take the largest connected component of this graph and make it undirected by ignoring edge directions as we are primarily interested in the topological structure rather than the specific flow direction of traffic. A crucial first step in our research is to empirically validate the scaling of these separators. We seek to determine if they scale near $\mathcal{O}(n^{1/3})$, as suggested by prior work [DSW16], or if an alternative scaling, like $\mathcal{O}(n^{1/2})$, merely appears smaller due to a low constant factor. Figure 4.1 plots the size of separators against the size of the corresponding subgraphs from which they are computed. Each data point (x, y) in this figure signifies that a subgraph containing x nodes possesses a separator of size y . The data points are generated by recursively applying nested dissection, computing separators first for the original graph and then for the subgraphs induced at each subsequent level.

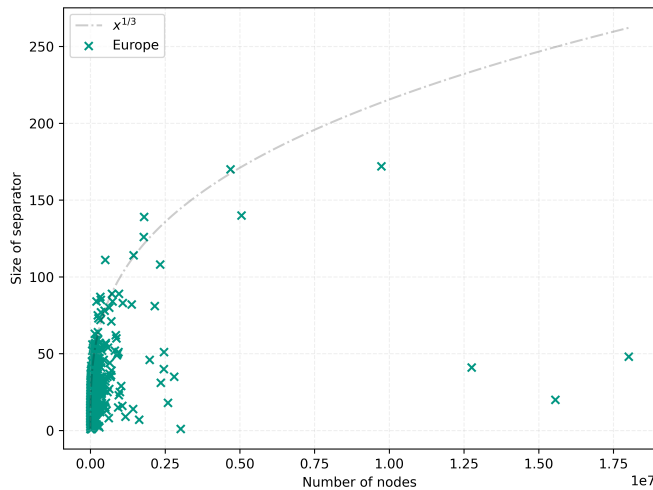


Figure 4.1: Empirical separator size versus subgraph size for the Europe road network. Each point represents a subgraph and its corresponding separator size.

Initial observations reveal outliers, particularly for very large subgraphs corresponding to continental or country scales. Specifically, analysis of the top-level separator structure for the Europe graph shows that the Scandinavian peninsula can be disconnected via separators significantly smaller than the general trend would suggest. This is due to specific geographic bottlenecks, as illustrated in Figure 4.2. Such outliers at the largest scales appear to be heavily influenced by macroscopic geographic features rather than intrinsic network structure representative of typical road networks. Consequently, these data points may not accurately reflect the general separator properties inherent in the finer structure of the road network graph. To mitigate the influence of these large-scale geographical artifacts and focus on more representative structural properties, our analysis primarily considers subgraphs with fewer than 10,000,000 nodes.

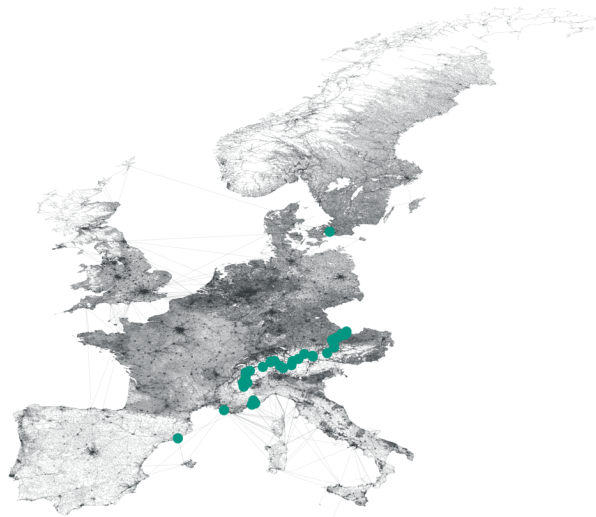


Figure 4.2: Illustration of a geographically influenced outlier at the continental scale: removal of a few nodes disconnects the entire Scandinavian peninsula.

For enhanced visibility, particularly concerning the numerous data points corresponding to smaller subgraphs, and to avoid overrepresentation of larger subgraphs, we will also present data a log-log scale. This logarithmic scaling offers the additional advantage that a polynomial relationship between separator size y and subgraph size x , such as $y \propto x^c$, manifests as a linear trend in the log-log plot, facilitating the identification of potential power-law dependencies. Furthermore, to improve the interpretability of the visualization and emphasize the underlying trend over individual fluctuations or outliers present at various scales, the data points are aggregated into bins. Let b be the number of bins chosen for the aggregation. Let x_{\max} denote the maximum observed subgraph size, assuming $x_{\max} > 0$. A data point (x, y) is assigned to the bin with index $\lfloor \frac{x \cdot b}{x_{\max}} \rfloor$. After assigning all points to their respective bins, a single representative point is computed for each non-empty bin. This representative point (\bar{x}_i, \bar{y}_i) for bin i is determined by calculating the arithmetic mean of the x coordinates and the arithmetic mean of the y coordinates of all data points (x, y) assigned to bin i . A primary consideration for using the arithmetic mean was the nature of subsequent analysis steps, such as curve fitting. While methods like box plots offer detailed distributional insights, they do not provide the single-point representation required for these analyses. Furthermore, the choice

of the mean over the median, which is known for its robustness to outliers, was deliberate in this context. Outliers within bins are not necessarily disregarded as noise but are considered potentially informative data points reflecting relevant variations. Supporting this decision are the mean's straightforward calculation, computational efficiency, and clear interpretation as the centroid or center of mass of the points within the bin.

When constructing the log-log plot, this binning procedure is applied to the logarithmically transformed data. Figure 4.3 illustrates the binned data points on logarithmic axes alongside the non-binned data.

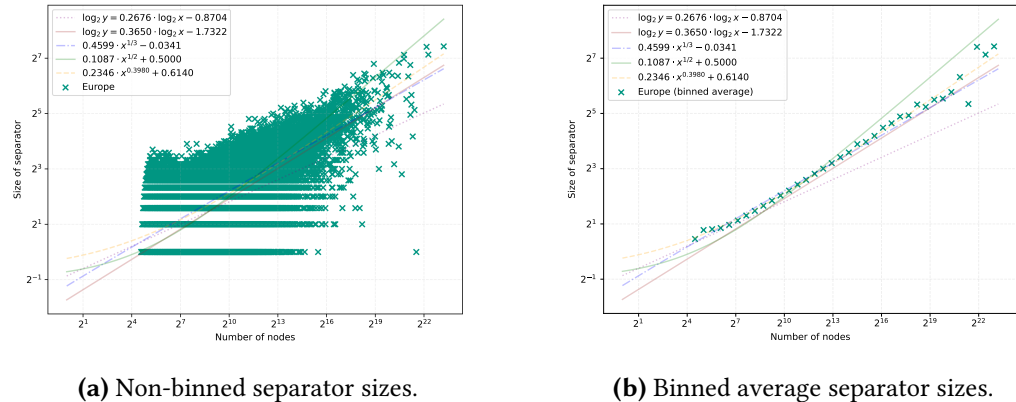


Figure 4.3: Separator sizes of Europe on logarithmic axes (subgraphs < 10M nodes).

To quantify the relationship between separator size y and subgraph size x , we perform statistical curve fitting on the empirical data. A preliminary linear regression applied to the logarithmically transformed data across all points yields a fitted line $\log y \approx 0.2676 \log x - 0.8704$. This initial fit suggests a scaling behavior close to $\mathcal{O}(x^{0.2676})$, seemingly better than the hypothesized $\mathcal{O}(x^{1/3})$. However, this result is likely skewed by the large number of small subgraphs, which may not fully represent the scaling trend at larger sizes. To test this hypothesis, we perform a second regression exclusively on data points corresponding to subgraphs with more than $2^8 = 256$ vertices. This analysis yields the relationship $\log y \approx 0.3650 \log x - 1.7322$.

We report these coefficients to four decimal places as is conventional in scientific reporting. However, we acknowledge that this level of precision might overstate the certainty of the fit, given the inherent noise in empirical measurements from complex graph algorithms. Minor variations in experimental conditions, such as changing the random seed used in the nested dissection algorithm, can cause slight fluctuations in these fitted parameters. Therefore, the reported values should be interpreted as indicative of the general trend rather than as exact constants.

We also explore direct non-linear curve fitting to the original data points using several functional forms. Fitting $y = a \cdot x^{1/3} + b$ results in $y \approx 0.4599 \cdot x^{1/3} - 0.0341$ with an R^2 value of 0.4724. Fitting $y = a \cdot x^{1/2} + b$ yields $y \approx 0.1087 \cdot x^{1/2} + 0.5000$, but with a very low R^2 value of 0.0873, suggesting that a square-root dependency is unlikely. A more general power-law fit $y = a \cdot x^c + b$ results in $y \approx 0.2346 \cdot x^{0.3980} + 0.6140$ with an R^2 value of 0.4832. While these direct fits indicate a scaling exponent potentially closer to 0.4 than to 1/3 or 1/2, the overall R^2 scores remain relatively low, indicating only a moderate goodness of fit. These fitted curves are visualized in Figure 4.3. The fitted lines displayed in Figure 4.3b are those computed

from the non-binned data and shown previously in Figure 4.3a. They are reproduced here to facilitate comparison with the binned averages, although visual alignment may differ due to the underlying averaging represented by the bins.

Observations from the binned log-log plot in Figure 4.3b suggest that the linear relationship between $\log y$ and $\log x$ is most consistent within a specific range of subgraph sizes. The data appear particularly stable for subgraph sizes x between approximately 2^7 and 2^{18} nodes. For subgraphs smaller than 2^7 nodes, a slight upward deviation is discernible, potentially reflecting behavior closer to grid-like structures at very small scales. For subgraphs larger than 2^{18} nodes, the increased scatter might be attributed to the limited number of data points available for aggregation in these higher size ranges.

To obtain a more robust estimate of the scaling exponent, we focus our analysis on the data within this more stable range ($2^7 \leq x \leq 2^{18}$). Furthermore, performing the linear regression on the binned data points offers two key advantages. Firstly, binning averages out the influence of individual outliers within each bin. Secondly, it gives more equal weight to different orders of magnitude in subgraph size, mitigating the numerical dominance of the numerous small subgraphs in the dataset.

Applying linear regression to the mean coordinates of the bins within the selected range on the log-log scale yields the fitted line:

$$\log y \approx 0.3702 \log x - 1.5512 \quad \Longleftrightarrow \quad y \approx 0.3411 \cdot x^{0.3702}$$

This fit demonstrates an exceptionally high coefficient of determination ($R^2 \approx 0.9994$), indicating that the linear model explains almost all the variance in the binned log-transformed data. The statistical significance of the slope is confirmed by an extremely low p-value ($p \approx 1.64 \times 10^{-20}$).

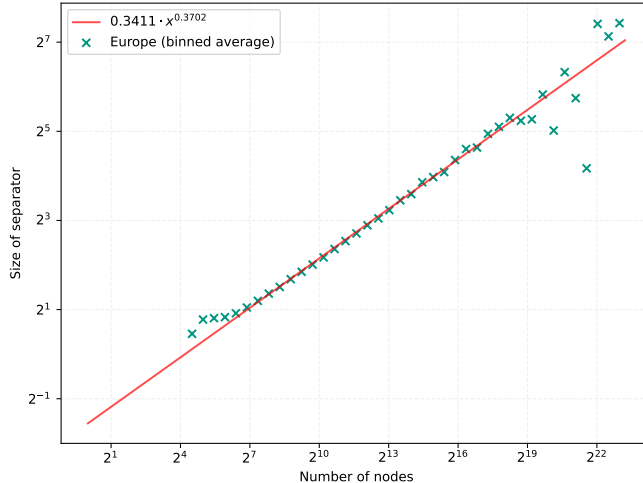


Figure 4.4: Linear regression fit to the binned data of separator size versus subgraph size, plotted on logarithmic axes. The regression considers only bins corresponding to subgraph sizes (x) in the range $2^7 \leq x \leq 2^{18}$.

Visually, this line provides an excellent fit to the binned data points within the considered range, as illustrated in Figure 4.4. The slope of 0.3702 in the log-log regression corresponds to the exponent in the power-law relationship $y \propto x^c$. Based on the high quality of the fit ($R^2 \approx 0.9994$) and the statistical significance of the result, we can state with high confidence that the observed scaling behavior is close to, yet slightly above, $\mathcal{O}(n^{1/3})$.

Initial Deviations in Separator Scaling Empirical studies of road networks reveal a notable deviation in separator scaling for small subgraphs. This deviation is particularly apparent for subgraphs containing approximately 2^6 vertices, where separator sizes are often larger than the general scaling trend would suggest, as can be observed in Figure 4.3a. Figure 4.5 presents a histogram that further visualizes the distribution of these separator sizes. This phenomenon corresponds with the observation of a higher meshedness coefficient for smaller, denser urban cores, as discussed in Section 4.2. The more grid-like, densely connected structure implied by a higher meshedness may contribute to these comparatively larger separators at smaller scales.

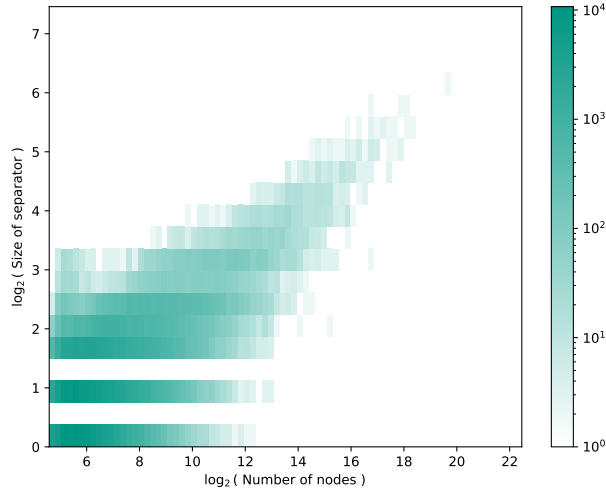


Figure 4.5: Histogram illustrating the distribution of separator sizes for road networks. A slight increase in relative separator size is observed for graphs with approximately 2^6 nodes.

4.2 Degree Distribution

Road networks are characteristically sparse graphs. The DIMACS Europe road network dataset from PTV [PTV09], used in our experiments, exemplifies this with an average vertex degree of approximately 2.5. Figure 4.6 presents this network's degree distribution, which reveals that a vast majority of vertices (approximately 99.8%) have a degree less than 5. The maximum degree observed in this dataset is 12, attained by a single node.

Implications for Other Structural Metrics The degree distribution and overall sparsity also influence other structural metrics, particularly when considering the near-planar nature of road networks. One such metric for planar-like graphs is the meshedness coefficient, α ,

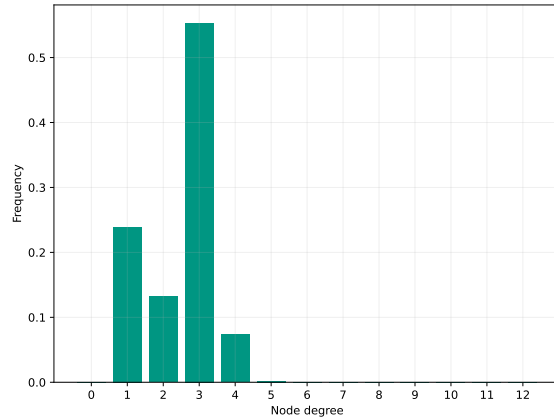


Figure 4.6: Degree distribution of the DIMACS Europe road network [PTV09]. The x-axis represents vertex degree, and the y-axis indicates the fraction of vertices.

which quantifies the density of cycles or bounded faces within a planar graph [Buh+06]. It is calculated as the ratio of a graph's actual number of bounded faces ($m - n + 1$) to the maximum possible for a planar graph with n vertices and m edges, via the formula $\alpha = \frac{m-n+1}{2n-5}$. This coefficient ranges from 0 for trees to 1 for maximal planar graphs. Buhl et al. suggest that this coefficient can also be used to gauge a network's robustness to disconnections and its cost in terms of total edge length [Buh+06]. For the DIMACS Europe road network, after applying a planarization procedure (detailed in Section 4.3), we compute a meshedness coefficient $\alpha \approx 0.1166$. This relatively low value underscores the network's general sparsity and somewhat tree-like macroscopic structure. For context, the meshedness coefficient for a square grid approaches 0.5 as $n \rightarrow \infty$, while a full Delaunay triangulation, as an almost maximal planar graph, approaches 1. In contrast to the continental scale, urban cores are typically more densely meshed. To illustrate this, we analyze the inner-city structure of Karlsruhe, a sub-region available within the PTV/DIMACS dataset. Representations of roads often include long chains of degree-2 vertices that model segments rather than structurally significant junctions. Such modeling can artificially skew the meshedness coefficient to lower values because adding these degree-2 vertices increases n without increasing the graph's cycle count ($m - n$). Therefore, to analyze the underlying junction-based topology independent of this modeling artifact, contracting degree-2 vertices is a valuable normalization step. After applying this contraction, the graph for Karlsruhe's inner city yields a meshedness coefficient of approximately 0.25. This higher value reflects the more grid-like layout characteristic of dense urban centers. Applying this same normalization step to the entire DIMACS Europe graph has a less pronounced impact, increasing its meshedness coefficient only slightly from 0.1166 to 0.1317.

4.3 Planarity

Road networks can be modeled as nearly planar graphs, meaning they permit an embedding in the plane with a limited number of edge crossings. Empirical evidence suggests that the number of such crossings in real-world road networks is typically on the order of $\mathcal{O}(n)$, a

sub-linear count relative to the number of edges [EG08]. It is a well-known result in graph theory that planar graphs admit $\frac{2}{3}$ -balanced separators of size $\mathcal{O}(n^{1/2})$ [LT79]. A relevant inquiry is whether the near-planarity of road networks is a critical feature that influences their separator properties, or if the occasional non-planar elements are merely incidental. This prompts the question of how separator sizes are affected when road networks are transformed into strictly planar graphs.

To obtain a planar representation, we begin with the existing graph structure where vertices possess associated geometric coordinates. Each edge is interpreted as the straight line segment connecting the coordinates of its incident vertices. The algorithm then identifies all geometric intersection points between these line segments. A new vertex is introduced into the graph at the coordinates of each detected intersection, provided this point does not coincide with an existing vertex. Any original edge containing one or more such intersection points is then removed and replaced by a sequence of new, shorter edges that connect the original endpoints and the new intersection vertices in their linear order. This process transforms the initial graph into a planar graph embedding by explicitly representing all edge crossings as vertices. For efficient execution, we utilize a spatial index over the bounding boxes of all edges. This structure enables rapid identification of potential intersections by querying for overlapping bounding boxes, which can then be verified for actual crossings. While other approaches exist, such as the Bentley-Ottmann algorithm for general line segment intersection [BO79] or linear-time algorithms tailored for graphs with a sublinear number of crossings [EGS10], our spatial index-based method is chosen for its implementation simplicity, as performance is not a critical concern for this pre-processing step. Since a single edge may cross multiple other edges, intersection points are sorted along each original edge before new sub-edges are introduced. Pseudo-code for this planarization algorithm is provided in Algorithm 4.1.

Algorithm 4.1: Simple planarization algorithm

Input: Non-planar graph $G = (V, E, pos)$.

Output: Planarized version of G .

```

1 spatial_index ← load(bounding_boxes( $E$ ))
2 crossings ← {}
3 forall  $e$  in  $E$  do
4   forall candidates  $c$  in spatial_index.query( $e$ ) do
5     if  $c$  intersects  $e$  then
6       crossings[ $e$ ].append( $c$ )
7       crossings[ $c$ ].append( $e$ )
8   forall ( $e$ , crossed) in crossings do
9      $G$ .remove( $e$ )
10    vertices ← get_intersection_vertices( $e$ , crossed)
11    sort vertices along  $e$ 
12    add_new_edges( $e$ , vertices)

```

We apply this planarization method to several real-world road networks. The Karlsruhe network, with approximately 120,000 nodes, has around 2,500 crossings. The Germany network, comprising about 6 million nodes, has approximately 100,000 crossings. Finally, the Europe network, with around 18 million nodes, contains about 300,000 crossings. These numbers are a little higher than the $\approx \mathcal{O}(n^{1/2})$ intersection counts reported in some prior

studies, but are of a comparable order of magnitude [EG08]. The differences could be explained by our modeling of edges as straight lines rather than more complex curves, and might be mitigated by using a more detailed road network model like OpenStreetMap.

Analysis of separator sizes shows minimal variation post-planarization. We identify $\frac{2}{3}$ -balanced separators with sizes still scaling approximately as $\mathcal{O}(n^{1/3})$, aligning with the values from the original non-planar graphs. A comparison of the separator sizes in the planar and non-planar versions of the Europe network is depicted in Figure 4.7.

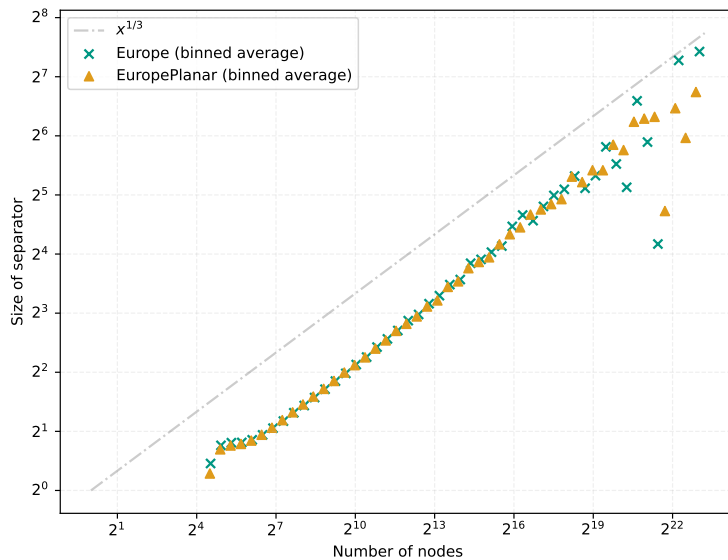


Figure 4.7: Comparison of separator sizes in the European road network: planar vs. non-planar versions.

Our findings indicate that separators in non-planar road networks closely resemble those in their planarized counterparts in terms of overall scaling. However, a separator from the original graph is not always directly valid in the planarized version, as edge crossings can create new paths between previously separated components. We investigated this by traversing a single arm of a nested dissection and checking if the separators for the original subgraphs remain valid in their planarized versions. Approximately one-third of these separators can be applied directly without modification, with this being more probable for smaller subgraphs that are less likely to contain edge crossings. When a separator is invalidated by the new topology, different outcomes are possible. In many cases, an original separator must be augmented with additional nodes to restore the partition, resulting in a slightly larger separator. Figure 4.8 provides a real-world example, visualizing how a separator from the non-planar Karlsruhe network has to be augmented to remain valid after planarization.

The new structure created by planarization can also reveal a new, more efficient separator that is smaller than the original, as shown in the example in Figure 4.9. Conversely, there are also rare cases in which no better separator can be found in the planarized graph, as illustrated in Figure 4.10.

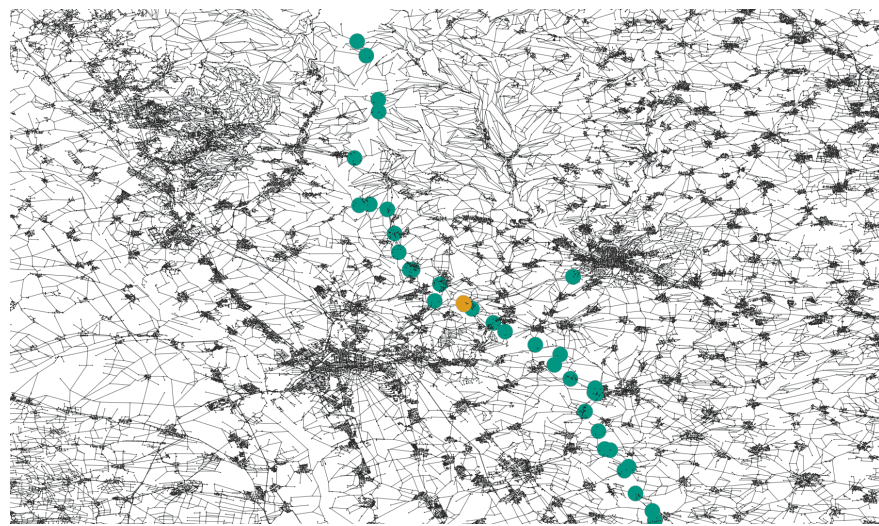
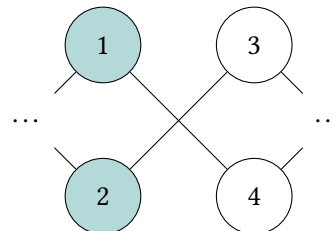
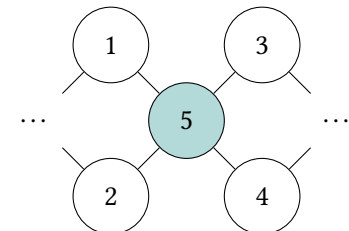


Figure 4.8: Visualization of a top-level separator for the road network of Karlsruhe. Vertices colored teal represent the separator nodes identified within the original, non-planar graph. Orange vertices indicate the additional nodes required to establish a valid separator for the planarized version of the network. The single teal vertex on the right is a highway intersection whose inclusion is necessary for the separator.

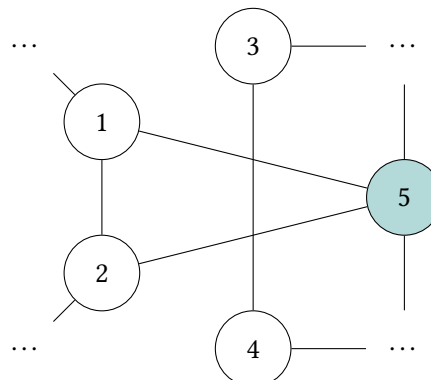


(a) Separator in non-planar graph.

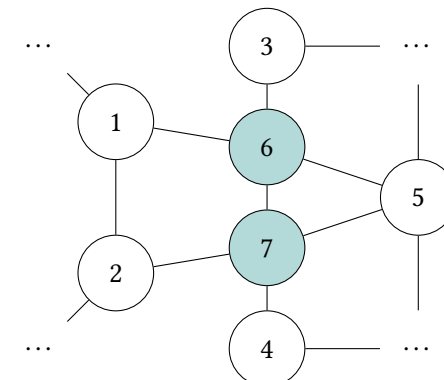


(b) Separator in planarized graph.

Figure 4.9: Example of a separator that decreases in size after planarization.



(a) Separator in non-planar graph.



(b) Separator in planarized graph.

Figure 4.10: Example of a separator that increases in size after planarization.

These findings highlight that the near-planar structure of road networks has minimal impact on separator size, suggesting that such networks can typically be analyzed as planar graphs without loss of essential separator properties.

4.4 Hierarchy

Real-world road networks possess a hierarchical structure. Different types of roads cater to different travel distances and volumes, forming levels within the network. Depending on the specific taxonomy used, road networks are categorized into various numbers of classes or levels. For instance, a common classification for Germany includes:

- Federal Motorways (Bundesautobahnen)
- Federal Highways (Bundesstraßen)
- State Roads (Landesstraßen)
- District Roads (Kreisstraßen)
- Municipal Roads (Gemeindestraßen)

4.5 Diameter and Distance Distributions

The diameter of a graph, defined as the longest shortest path between any pair of vertices, is a fundamental metric characterizing its overall extent and the efficiency of traversal. In our analysis, we focus specifically on the hop diameter, where each edge has a uniform weight of 1. This choice is motivated by the desire to understand the graph's topological and hierarchical structure. To ensure that the measured hop distances are structurally meaningful, we first pre-process all graphs and subgraphs by contracting vertices of degree 2. This effectively treats long chains of such vertices as single conceptual edges.

Computing the exact diameter of a general graph can be computationally intensive. A simple two-step Breadth-First Search approach, performing a BFS from a random node to find the furthest vertex and then a second BFS from that result, is only guaranteed to find the exact diameter in certain graph classes, such as trees, and provides at best a 2-approximation for general graphs. Therefore, to obtain exact diameter values, we employ the more sophisticated iFUB (iterative Fringe Upper Bound) algorithm as described by Crescenzi et al. [Cre+13]. This algorithm successfully computes the exact graph diameter in reasonable time on road networks without resorting to a full all-pairs shortest path calculation.

We investigate how the hop diameter scales with graph size by analyzing subgraphs obtained from the nested dissection process. Our empirical analysis of these contracted subgraphs indicates that their diameter scales approximately as $\mathcal{O}(n^{1/3})$. This observed scaling behavior is illustrated in Figure 4.11.

Beyond the maximum path length (diameter), we also analyze the complete distributions of both weighted distances and hop counts to understand typical path lengths. Computing all-pairs shortest paths is computationally prohibitive for large graphs. Therefore, we approximate these distributions by sampling 10^4 source nodes and performing a one-to-all shortest path query from each. For the hop distance calculation, the graph is pre-processed by contracting all degree-2 vertices, consistent with our diameter analysis. Figure 4.12 visualizes the resulting probability density functions and cumulative distribution functions for both metrics. Both

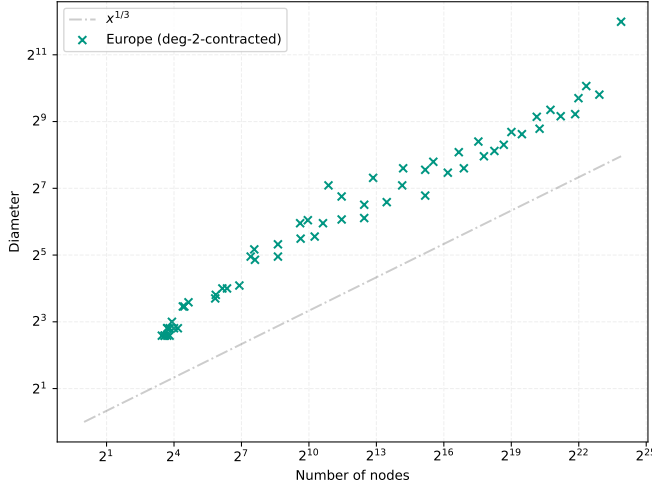
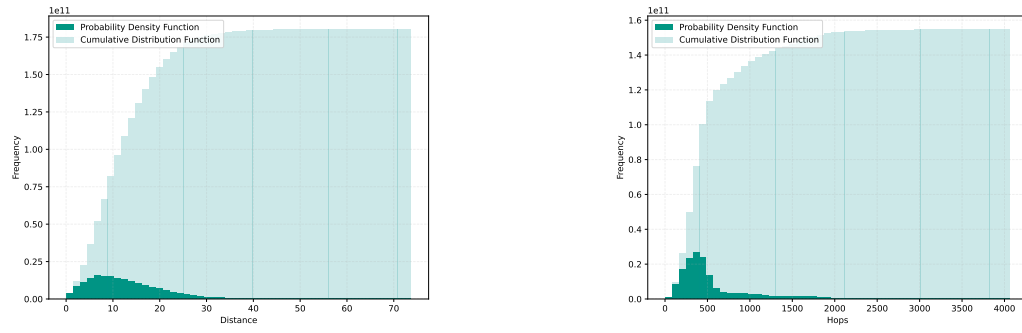


Figure 4.11: Empirical scaling of hop diameter with the number of vertices n for nested dissection subgraphs of road networks (with degree-2 nodes contracted). The observed scaling is approximately $\mathcal{O}(n^{1/3})$.

distributions are strongly left-skewed, indicating that most pairs of vertices are relatively close, followed by a long tail of outliers that are hard to reach. The hop distribution (Figure 4.12b) is more sharply skewed with a steeper drop-off compared to the weighted distance distribution (Figure 4.12a). This difference may indicate a functional hierarchy within the network. The steep hop distribution could reflect a backbone of “highway” edges that efficiently connect distant regions in a small number of hops, while the slower decay in the weighted distance distribution accounts for the actual, larger geographic distances covered.



(a) Distribution of weighted path distances.

(b) Distribution of path lengths in hops.

Figure 4.12: Probability density functions (PDFs) and cumulative distribution functions (CDFs) for weighted distances and hop counts in the PTV Europe road network, approximated from 10,000 one-to-all queries.

5 Synthetic Graph Generation for Feature Isolation

To better understand the underlying reasons for the small separators observed in road networks, we investigate the influence of specific graph properties in isolation. This chapter details our approach to generating synthetic graph classes that exhibit selected features characteristic of road networks, such as low average degree, specific degree distributions, or properties related to planarity and locality. By analyzing the separator sizes within these synthetic graphs, we aim to determine which properties, or combinations thereof, are crucial for enabling small separators.

5.1 Degree Distribution

Our initial focus is on isolating the effect of the degree distribution. To examine whether a low average degree is sufficient to yield small separators, we generate connected random graphs matching this average degree. The generation process involves two main steps. First, a random spanning tree is created for a given set of n vertices using the algorithm described by Broder [Bro89]. This algorithm performs a random walk starting from an arbitrary vertex on the complete graph of n vertices. An edge becomes part of the spanning tree the first time a vertex is discovered via that edge during the walk. The process continues until all vertices are visited, resulting in a uniformly sampled random spanning tree in expected time $\mathcal{O}(n \log n)$.

It is noteworthy that random trees generated in this manner exhibit properties distinct from those of road networks. For instance, the diameter of such random trees is known to be in $\mathcal{O}(n^{1/2})$ [CK87]. This differs from our empirical observations on road networks. This observed diameter growth of these trees is visualized in Figure 5.1.

Following the generation of the initial random spanning tree, we proceed to the second step: adding edges randomly between pairs of non-adjacent vertices. This edge addition continues until the target average degree of 2.5 is reached for the entire graph. The resulting graphs, by construction, lack the inherent locality often present in road networks. A consequence of adding these random edges is a notable decrease in the graph diameter. For example, graphs generated with one million nodes using this method exhibit diameters of approximately 40.

These synthetic graphs do not replicate the small separator sizes observed in real-world road networks. Experiments yielded large top-level separators, whose sizes scaled approximately linearly with the graph size n , i.e., as $\mathcal{O}(n)$. However, separators found during subsequent recursive partitioning behaved differently: their sizes were significantly smaller than this linear trend would predict for the corresponding subgraph sizes. We attribute this deviation to structural changes in the subgraphs induced by the partitioning process. Specifically, the large top-level separators often remove a significant fraction of the non-tree edges that were originally added to achieve the target average degree of approximately 2.5 in the parent graph. Consequently, the resulting subgraphs become sparser, our observations showed a decrease in average degree to approximately 2.2, and increasingly dominated by their underlying tree

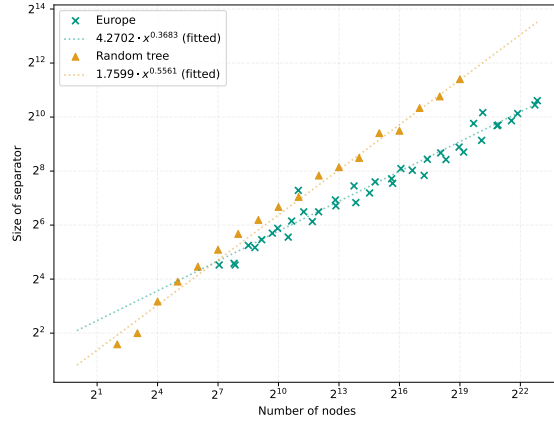


Figure 5.1: The plot shows the diameter (y-axis) as a function of the number of nodes n (x-axis, log scale).

structure. This shift means these subgraphs effectively belong to a different, more tree-like graph class than initially generated, which naturally leads to smaller relative separator sizes. The separator scaling is illustrated in Figure 5.2.

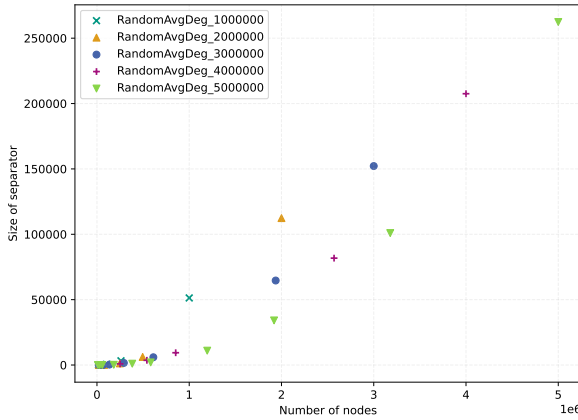


Figure 5.2: Separator sizes observed in random graphs generated with an average degree matching the DIMACS Europe road network (≈ 2.5).

We also conducted experiments to generate graphs that match not just the average degree but also the specific degree distribution of a reference road network. This process involved starting with a random tree and then adding edges subject to constraints: an edge (u, v) was incorporated only if doing so would not cause either vertex u or v to exceed the degree count specified for them by the target distribution, which was sampled from the road network. The separator results from this more refined approach were largely similar to those from graphs matching only the average degree.

It is important to note, however, that precisely replicating a target degree distribution with this method is not possible. While the maximum degree in the generated graphs generally aligns with that of the reference network, discrepancies emerge, particularly for nodes with higher degrees. The underlying random spanning tree structure tends to produce a higher proportion of such nodes compared to actual road networks. For instance, in synthetic graphs designed to match the Europe dataset [PTV09], approximately 1.8% of vertices had a degree of 5 or greater, exceeding the 0.2% observed in the real dataset. Thus, our process achieves an approximation of the target distribution. Nevertheless, we posit that this approximation is sufficiently close for our investigation, as the relatively small fraction of vertices with degrees deviating from the target constraints is unlikely to fundamentally alter the observed separator scaling behavior for this graph class.

These findings reinforce the suggestion that the specific degree distribution, in isolation, is likely insufficient to fully explain the empirically observed small separators. Nevertheless, the low average degree remains a noteworthy property. While a lower average degree may not fundamentally alter the inherent asymptotic separator scaling characteristic of a specific graph class, it often correlates with smaller constant factors in the observed separator sizes; the practical effect of sparsity on separator magnitudes will be discussed further in a subsequent section.

Beyond the influence of average degree on constant factors, it is also crucial to recognize that neither a low average degree nor a specific sparse degree distribution are strict prerequisites for achieving small separators. Indeed, graphs can be constructed that possess both deliberately small separators (as a function of n) and a high average degree. Consider, for instance, the following recursive graph generation scheme designed to produce a graph on n vertices with a target separator of size $s(n)$ (e.g., $s(n) = n^{1/3}$): First, a separator clique S on $s(n)$ vertices, denoted $K_{s(n)}$, is designated. Then, two subgraphs, G_1 and G_2 , each containing approximately $(n - s(n))/2$ vertices, are generated recursively by the same procedure (with a base case, such as returning a clique K_n , for small n , e.g., when $n \leq 2$). Finally, every vertex in G_1 is connected to every vertex in S , and likewise, every vertex in G_2 is connected to every vertex in S . By construction, the set S (forming a $K_{s(n)}$) is a separator of size $s(n)$ that partitions the graph into G_1 and G_2 . Despite this small, built-in separator, the average degree of the resulting graph can be substantial. For example, using $s(n) = n^{1/3}$, a graph of $n = 10^4$ vertices generated via this method has an average degree of approximately 169 and a maximum degree of 6202. This example illustrates that specific structural properties enabling small separators can coexist with high overall edge density, suggesting that the global organization of edges, beyond mere sparsity, significantly influences separator characteristics.

5.2 Locality

Our investigation into locality begins by exploring how “closeness” defined by an existing network structure, rather than by direct geometric proximity in an embedding, influences graph properties.

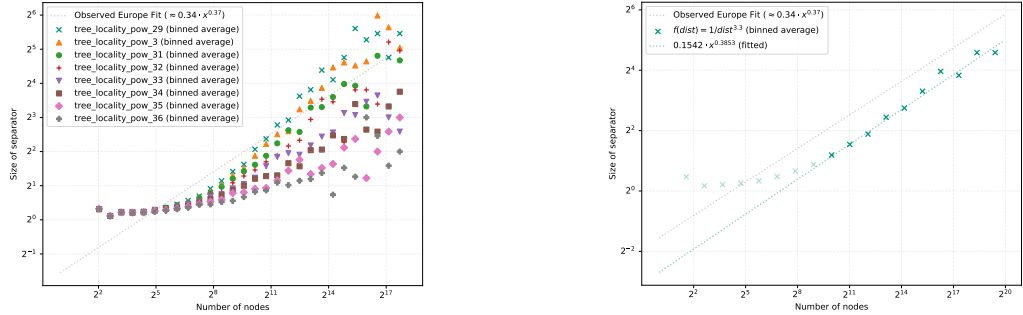
Tree Locality For this first approach, we simulate locality by considering distances within an initial random spanning tree, which serves as a foundational sparse connected graph. The premise is that vertices closer to each other along paths in this underlying tree are more likely candidates for new, direct connections, potentially reflecting how topological proximity in an existing network might drive the addition of new links. To implement this, we

begin by generating a random spanning tree using Broder’s algorithm [Bro89]. Subsequently, edges are added between non-adjacent vertices until a target average degree, chosen as 2.5 to approximate that of road networks, is achieved. To incorporate this form of locality, the probability of adding an edge between two vertices u and v is made dependent on their distance $\text{dist}_T(u, v)$ within this initial spanning tree T . Specifically, we select a random vertex x and choose a second vertex y with a probability related to $f(\text{dist}_T(x, y))$, where f is a decreasing function.

The computation of tree distances $\text{dist}_T(x, y)$ for numerous candidate pairs (x, y) is a critical step in this edge addition phase. Initially, for each randomly selected vertex x and potential neighbor y , a Breadth-First Search (BFS) is performed within the tree T to find $\text{dist}_T(x, y)$. While BFS on a tree is linear in the number of vertices n , repeated invocations for many pairs prove computationally intensive for larger graphs. To accelerate this, we adopt an approach based on Lowest Common Ancestor (LCA) queries. This involves an initial precomputation of an LCA data structure for the tree T , which can be achieved in $\mathcal{O}(n \log n)$ time (e.g., using an Euler tour and sparse table for Range Minimum Queries, enabling $\mathcal{O}(1)$ query time). Once the LCA of two nodes u and v is known, their tree distance is given by $\text{dist}_T(u, v) = \text{depth}(u) + \text{depth}(v) - 2 \cdot \text{depth}(\text{lca}(u, v))$. Although a single BFS is also linear, this LCA-based method yields a significant speedup, due to improved parallelization opportunities for the overall process. Nevertheless, even with this optimization, the approach of potentially considering many y candidates for each x (full sampling) remains slow for large graphs. We also test capping the BFS search for y (e.g., after a fixed number of hops or visited nodes, such as 50,000) as a potential optimization. However, this leads to an artificial steep cutoff in the observed separator characteristics once graph or component sizes effectively exceed the scale imposed by this search limit.

Experiments exploring various decay functions f yield diverse results. Simple functions like $f(\text{dist}) = 1/\text{dist}$ result in graphs that still exhibit large top-level separators scaling as $\mathcal{O}(n)$, although subgraphs from nested dissection show the previously noted super-linear decrease in separator size. Conversely, rapidly decaying functions like $f(\text{dist}) = 2^{-\text{dist}}$ produce graphs with separators of almost constant size, likely because the graph structure remains very close to that of the initial tree, with insufficient long-range connections. Through further experimentation and fine-tuning, it is found that a function of the form $f(\text{dist}) = 1/\text{dist}^{3.3}$ is able to produce separator sizes that closely approximate the desired $\mathcal{O}(n^{0.37})$ scaling (see Figure 5.3b). Figure 5.3 provides a visual comparison of the separator sizes for various decay functions.

Despite achieving the target scaling with $f(\text{dist}) = 1/\text{dist}^{3.3}$, this result offers limited insight into the intrinsic structural properties of real road networks that lead to their small separators. The success with this specific, empirically derived exponent appears to be more a consequence of parameter tuning within this particular generative model (a random tree backbone augmented by edges based on tree distance) rather than an elucidation of fundamental graph characteristics. While the general concept of incorporating locality based on distances within an existing network structure remains potentially insightful, as such a metric might capture aspects beyond mere Euclidean distance, like travel time reduction incentives, our preliminary results with simpler functions indicate a high sensitivity to the choice of f . Identifying a theoretically grounded or easily interpretable function f that produces $\mathcal{O}(n^{0.37})$ scaling proves non-trivial. Given these considerations and the decision to prioritize other structural properties like geometric locality, planarity, and explicit hierarchy, we opt not to pursue an exhaustive search for an optimal or more interpretable tree-distance decay function within the scope of this work.



(a) Separator scaling for various decay functions $f(\text{dist})$.

(b) Separator scaling for $f(\text{dist}) = 1/\text{dist}^{3.3}$ with line fit.

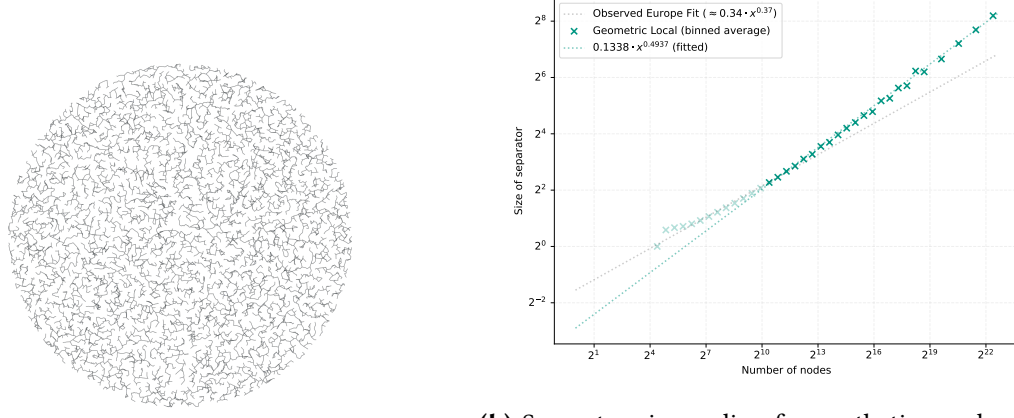
Figure 5.3: Separator size scaling for graphs generated with tree-distance based locality, comparing various decay functions and highlighting the fit for $f(\text{dist}) = 1/\text{dist}^{3.3}$.

Achieving the target $\mathcal{O}(n^{1/3})$ scaling with an empirically derived function like $f(\text{dist}) = 1/\text{dist}^{3.3}$ appears to be an outcome of model-specific parameter tuning rather than a revelation of intrinsic road network properties responsible for their small separators.

Geometric Locality Our second approach incorporates geometric locality directly. The core idea is to construct a graph starting with a basic connected structure and then augment it by adding only edges that connect geometrically close vertices, until a target average degree (e.g., 2.5) is achieved. To establish a meaningful threshold for 'closeness', we relate it to the natural scale derived from the spatial distribution of points. Specifically, we first sample n points uniformly within a defined spatial domain (e.g., a circle). We then compute the Minimum Spanning Tree (MST) of these points using Euclidean distances, via Kruskal's algorithm [Kru56]. The maximum edge length found within this MST, denoted ℓ_{\max} , serves as our threshold distance, capturing a characteristic length scale of the initial sparse connection of the points. The graph generation then proceeds by initially taking the edges of the MST. Subsequently, additional edges (u, v) between non-adjacent vertices are iteratively added, but crucially, only if their Euclidean distance is less than or equal to the threshold ℓ_{\max} . This edge addition process continues until the overall graph reaches the target average degree of 2.5. For efficient implementation of the edge addition step, we utilize spatial queries: for a randomly selected vertex u , we query for potential neighbors v within the radius ℓ_{\max} and randomly select one to connect to, avoiding multi-edges and self-loops.

Despite incorporating this explicit geometric constraint, the resulting synthetic graphs fail to exhibit the desired small separator sizes characteristic of road networks. To analyze the separator properties of these graphs, we computed recursive separators. Implementations from both IntertialFlowCutter and KaHIP were utilized for this purpose. Our experiments using these methods indicate that separators in these geometrically generated graphs scale approximately as $\mathcal{O}(n^{1/2})$. This outcome is visualized in Figure 5.4.

An alternative strategy we investigate for incorporating geometric locality is the construction of k-Nearest Neighbor (k-NN) graphs. In this model, each vertex connects via an edge to its k geographically closest neighbors, with proximity defined by the Euclidean distance between their embedded point coordinates. This approach does not utilize an underlying tree structure like some previously discussed methods. Connectivity in a k-NN graph is not



(a) Visualization of a graph with 10^4 nodes generated using the geometric locality method.

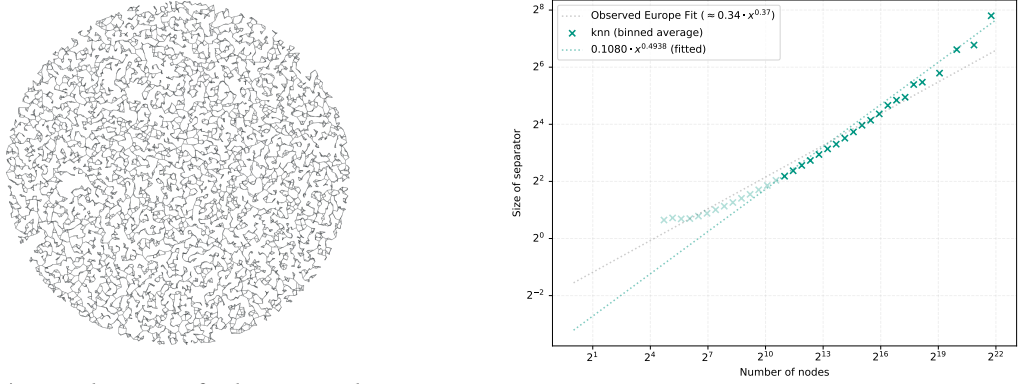
(b) Separator size scaling for synthetic graphs generated with geometric locality. The fit shown excludes subgraphs with ≤ 1000 vertices due to their disproportionately large separators.

Figure 5.4: Synthetic graph generation using geometric locality and analysis of separator sizes.

guaranteed; however, the size of the largest connected component generally increases with the value of k . For instance, with uniformly sampled points, a k as small as 3 often yields a primary connected component encompassing approximately 98% of all vertices. The average degree of the resulting graph is directly influenced by k . For point sets with non-uniform distributions, a larger k is typically required to achieve a similar level of overall connectivity. Our objective is to identify the smallest k that results in a largely connected graph, while also aiming for a low average degree, comparable to those of sparse road networks. The construction of k -NN graphs can be performed efficiently: following an initial $\mathcal{O}(n \log n)$ precomputation to build a spatial index such as a k -d tree, identifying the k nearest neighbors for each of the n vertices takes approximately $\mathcal{O}(k \log n)$ time per vertex, resulting in a total construction time of roughly $\mathcal{O}(nk \log n)$. Our experiments show that k -NN graphs generated in this manner also exhibit separator sizes that scale approximately as $\mathcal{O}(n^{1/2})$ as seen in Figure 5.5.

The results from the geometric locality approach suggest that merely combining low average degree with this specific form of locality is insufficient to reproduce the cubic root separator sizes of road networks.

Real-World Implications It is worth considering the practical implications of different asymptotic growth rates for separator sizes within the typical scale of road networks. While $\mathcal{O}(c_1 \cdot n^{1/3})$ and $\mathcal{O}(c_2 \cdot n^{1/2})$, with $c_2 \ll c_1$, diverge for large n , the actual separator sizes for graphs up to around 20 million vertices are similar. The performance of algorithms leveraging graph separators in real-world applications may be more influenced by the absolute sizes of separators achievable in practice, compared to a specific scaling model. Performance gains, for example for CCH [DSW16], could potentially be realized even if the separators behave like $c \cdot n^{1/2}$ for a sufficiently small c , as the absolute separator sizes remain manageable for networks of practical relevance.



(a) Visualization of a k-NN graph (for $k = 3$) generated from $n \approx 10^4$ uniformly sampled points.

(b) Separator size scaling for k-NN graphs $k = 3$, showing approximately $\mathcal{O}(n^{1/2})$ behavior.

Figure 5.5: Analysis of k-Nearest Neighbor (k-NN) graph of uniformly sampled points and separator scaling.

Initial Deviations in Separator Scaling Our synthetic graphs generated using the geometric locality approach exhibit a notable initial deviation in their separator scaling at small graph sizes. As visualized in the histogram in Figure 5.6, there is an initial peak in relative separator size for small subgraphs, which then decreases before the data align with the more dominant scaling trend observed for larger graphs. This behavior is similar to the initial scaling deviations present in real road networks. This correspondence is noteworthy, as it suggests that our geometric locality model, despite failing to replicate the overall asymptotic separator scaling, may capture certain structural properties that are relevant at smaller graph scales and also present in actual road networks.

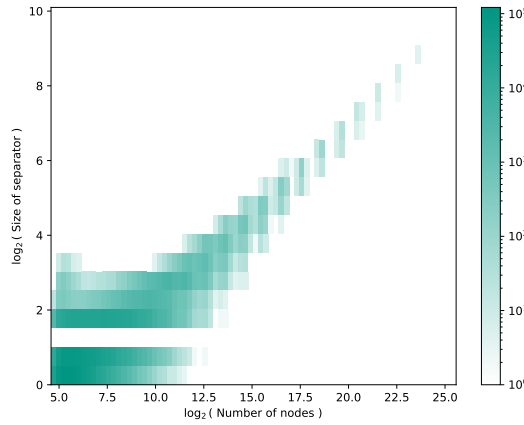


Figure 5.6: Histogram of separator sizes for synthetic graphs using geometric locality, illustrating an initial peak in relative separator size for small subgraphs (around 2^6 nodes).

Minimum Spanning Trees from Higher-Dimensional Embeddings Observing that real road networks often exhibit diameters smaller than the $\mathcal{O}(n^{1/2})$ found in our random tree models, we sought to determine if an initial tree structure with an intrinsically smaller diameter could lead to graphs with correspondingly smaller separators after edge augmentation. This motivated an exploration of generating initial trees as Minimum Spanning Trees (MSTs) of points sampled in d -dimensional Euclidean space, since the diameter of such an MST on n random points in d -space (for $d \geq 2$) is typically $\mathcal{O}(n^{1/d})$. Our experiments focused on the case where $d = 3$, for which the MST diameter is approximately $\mathcal{O}(n^{1/3})$. This involved sampling n points uniformly in 3D space and computing their MST. When edges were subsequently added to such 3D-MSTs to achieve a target average degree, the resulting graphs exhibited separators that scaled as $\mathcal{O}(n^{2/3})$. This scaling aligns with a general geometric intuition: a d -dimensional hypervolume containing n points is naturally bisected by a $(d - 1)$ -dimensional separator. If the characteristic linear extent of the volume is proportional to $n^{1/d}$, the $(d - 1)$ -dimensional measure (e.g., area for $d = 3$, length for $d = 2$) of such a separator would be proportional to $(n^{1/d})^{d-1} = n^{(d-1)/d}$. For $d = 2$, this yields the familiar $\mathcal{O}(n^{1/2})$ scaling for planar-like separators, and for our $d = 3$ experiment, it corresponds to the observed $\mathcal{O}(n^{2/3})$. While this suggests a general hypothesis that augmenting MSTs from d -dimensional point sets might lead to graphs with $\mathcal{O}(n^{(d-1)/d})$ separators, this avenue was not pursued further. Since for $d = 3$ the $\mathcal{O}(n^{2/3})$ separators do not offer an improvement over the $\mathcal{O}(n^{1/2})$ separators of planar graphs in the context of finding exceptionally small separators (and higher dimensions $d > 3$ would yield even larger exponents as $(d - 1)/d \rightarrow 1$), this specific line of inquiry was not extended.

5.3 Planarity

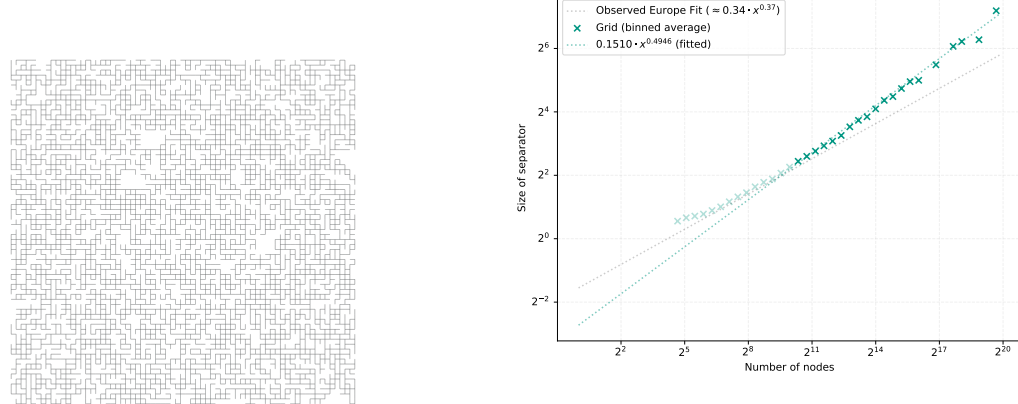
We examine separator properties in two classes of planar graphs: grids and Delaunay triangulations. These serve as fundamental theoretical models for planar structures relevant to the study of road networks due to their near-planarity. In our study, these graph classes are sparsified to achieve a low average degree (approximately 2.5), reflecting the sparsity observed in road networks. It is important to note that both of these models inherently introduce strong geometric locality, a property not required by all planar graphs. Our focus on these specific classes therefore introduces a bias towards investigating planar graphs that also possess this structural characteristic.

5.3.1 Grids

Our initial investigation focuses on grid graphs, a fundamental class known to possess separators scaling as $\mathcal{O}(n^{1/2})$. To align their sparsity with road networks, we generate modified grids with an average degree of approximately 2.5. The generation process starts with a square two-dimensional grid graph. Edges are then removed uniformly at random until the target average degree is reached over the entire graph. Subsequently, we identify and utilize the largest connected component for analysis. We observe that even without explicit mechanisms to prevent disconnection during edge removal, the largest connected component typically encompasses a large fraction of the initial vertices.

Analysis of these sparse grid graphs reveals separator sizes consistent with the $\mathcal{O}(n^{1/2})$ asymptotic behavior of complete grids. However, the constant factor associated with this scaling appears to be relatively small. Consequently, although the asymptotic limit behavior differs from the $\mathcal{O}(n^{1/3})$ scaling empirically observed for road networks, the absolute separator

sizes in these sparse grids are numerically similar to those of road networks for graphs up to typical sizes (e.g., around 20 million nodes). This finding highlights that sparsity, even within a simple planar structure like a grid, can lead to separators that are small in absolute terms for practical graph dimensions. Figure 5.7 illustrates a sample sparse grid and the observed separator scaling.

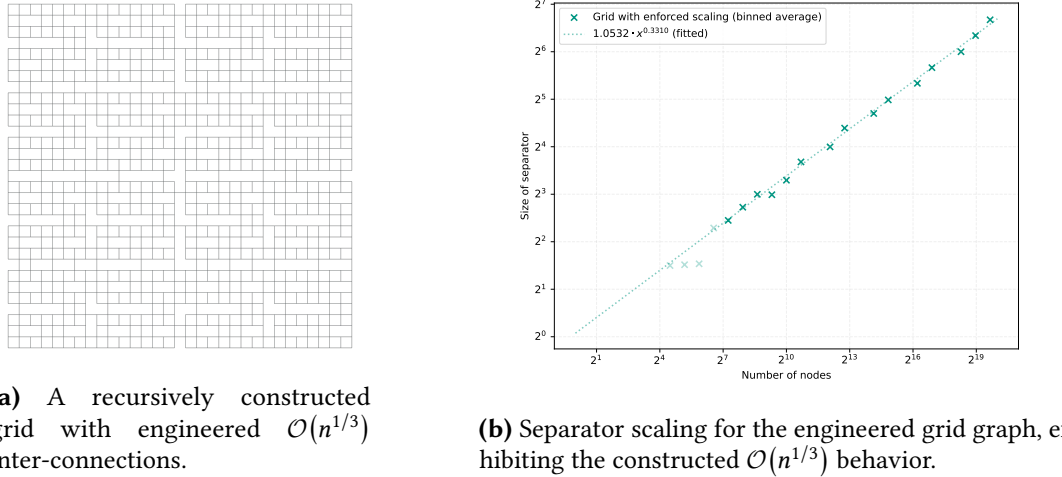


(a) Visualization of the largest connected component of a grid graph with $\approx 10K$ nodes after random edge removal to achieve an average degree of 2.5.

(b) Separator size scaling for sparse grid graphs (average degree 2.5). Separator sizes scale approximately as $\mathcal{O}(n^{1/2})$, but with a small constant factor leading to absolute sizes comparable to road networks at practical scales. The fit shown excludes subgraphs with ≤ 1000 vertices due to their disproportionately large separators.

Figure 5.7: Analysis of sparse grid graphs with average degree 2.5.

Analogous to the earlier example where dense graphs were constructed with a specific separator function, it is also possible to engineer grid-like graphs to exhibit a target separator scaling, such as $\mathcal{O}(n^{1/3})$. This approach involves a recursive construction: two smaller grid-like subgraphs, G_1 and G_2 , each comprising approximately $n/2$ vertices, are generated recursively. These subgraphs are then interconnected. Instead of forming dense connections or connecting all border vertices, a controlled number of edges, specifically chosen to match the desired separator size (e.g., approximately $n^{1/3}$ for the combined graph of size n), are added between G_1 and G_2 . For instance, this can be achieved by selecting approximately $n^{1/3}$ nodes along the boundary of one subgraph and connecting each to its closest corresponding node on the boundary of the other. The set of these inter-connecting edges (or one of their incident vertices) is, by construction, a separator of the desired $\mathcal{O}(n^{1/3})$ size. An example of such a graph and its resulting separator scaling is illustrated in Figure 5.8. While this method demonstrates that specific separator sizes can be achieved by deliberate construction in grid-like structures, it offers limited insight into which intrinsic graph properties of real road networks lead to their empirically observed small separators. The desired scaling is explicitly engineered into the generation process here, rather than emerging naturally from more fundamental characteristics.



(a) A recursively constructed grid with engineered $\mathcal{O}(n^{1/3})$ inter-connections.

(b) Separator scaling for the engineered grid graph, exhibiting the constructed $\mathcal{O}(n^{1/3})$ behavior.

Figure 5.8: Grid-like graph with engineered separators designed to scale as $\mathcal{O}(n^{1/3})$.

5.3.2 Delaunay Triangulations and Their Sparse Subgraphs

As a generalization of grid-like structures derived from point sets, Delaunay Triangulations (DT) are fundamental in computational geometry and serve as a basis for generating planar graphs. A DT for a given set P of discrete points in a plane is a specific triangulation such that for any triangle in $DT(P)$, its circumcircle (the unique circle passing through its three vertices) contains no other points from P in its interior. This “empty circumcircle” property ensures that DTs tend to maximize the minimum angle of all triangles in the triangulation, thereby avoiding overly elongated triangles. Our process for generating a baseline Delaunay graph involves sampling n points uniformly at random in a two-dimensional space and then computing their full Delaunay triangulation. Such a DT typically has an average vertex degree around 6. This observation is consistent with Euler’s formula for planar graphs ($|V| - |E| + |F| = 2$); for large triangulations, where most faces are triangles and each internal edge is shared by two faces ($3|F| \approx 2|E|$), the average degree $2|E|/|V|$ approaches 6.

Since an average degree of 6 is considerably denser than that of typical road networks (which is closer to 2.5), sparsification is necessary to create more comparable synthetic graphs. We explore several methods for this:

Random Edge Deletion One straightforward sparsification technique, similar to that applied in our grid experiments, involves randomly deleting edges from the full Delaunay triangulation until the target average degree of approximately 2.5 is achieved. The largest connected component of the resulting graph is then used for analysis.

Systematically Defined Subgraphs Alternatively, sparser graphs can be derived from the DT by retaining only those edges that satisfy stricter geometric conditions. Prominent examples of such Delaunay subgraphs include Gabriel Graphs (GG) [GS69] and Relative Neighborhood Graphs (RNG) [Tou80]. An edge (u, v) is part of a GG if the closed disk whose diameter is the segment uv contains no other point from the original set P . An edge (u, v) belongs to an RNG if no third point $w \in P$ is simultaneously closer to both u and v than u and v are to each other; that is, the lune formed by the intersection of two circles of radius

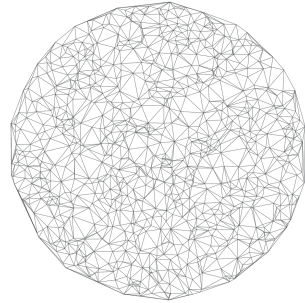
$\text{dist}(u, v)$ centered at u and v must be empty of other points. These definitions lead to the known hierarchical relationship: RNG \subseteq GG \subseteq DT. The RNG is particularly interesting from a network modeling perspective, as its construction criterion can be loosely interpreted in terms of economic viability for infrastructure: a direct road between two points u and v might not be constructed if an alternative path via a nearby third point w (e.g., $u \rightarrow w \rightarrow v$) exists without a significant detour. For uniformly sampled random points, the RNG yields an average degree of approximately 2.6 [Buh+06], which is close to the average degree of road networks.

Separator Analysis of Delaunay Variants We analyzed the separator properties of these different Delaunay-derived graphs: the full DT, the randomly sparsified DT (target average degree 2.5), the Gabriel Graph, and the Relative Neighborhood Graph, all generated from identical initial point sets to ensure comparability. Visualizations of these graph structures and their comparative separator scaling are presented in Figure 5.9. Despite the variations in density, ranging from an average degree of approximately 6 for the full DT down to about 2.6 for the RNG, our experiments indicate that the asymptotic separator scaling for all these Delaunay-derived planar graphs remains consistent with $\mathcal{O}(n^{1/2})$. The method of sparsification, whether through random edge deletion or by applying systematic geometric rules like those for GG or RNG, primarily influences the constant factors associated with the separator size. It does not, however, fundamentally alter the $\mathcal{O}(n^{1/2})$ scaling behavior tied to their underlying planar geometric nature. Even when the average degree very closely matches that of road networks (as in the RNG case), the separator scaling does not approach the $\approx \mathcal{O}(n^{0.37})$ observed empirically for road networks.

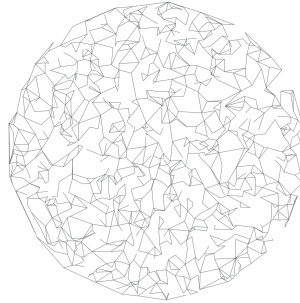
5.4 Highway Dimension

Given that the simpler graph models explored in previous sections do not reproduce the observed $\mathcal{O}(n^{1/3})$ separator scaling of road networks, we turn our attention to more complex generative processes designed to capture other relevant structural properties. One such property is highway dimension, introduced by Abraham et al. [AFGW10]. Intuitively, a graph possesses a small highway dimension if, for every radius $r > 0$, there exists a sparse set of vertices S_r such that every shortest path longer than r intersects S_r . A set is considered sparse if every ball of radius $\mathcal{O}(r)$ contains only a small number of vertices from S_r [AFGW10]. The significance of this property stems from the finding that low highway dimension provides provable performance guarantees for several important route planning algorithms, including REACH [GKW06], Contraction Hierarchies [GSSD08], Highway Hierarchies [SS05], Transit Node Routing [BFSS07], and SHARC [BD10].

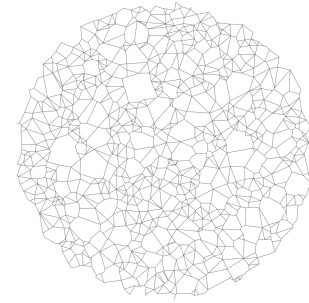
The work introducing highway dimension also proposes a synthetic graph generator (henceforth ABR generator) intended to produce graphs exhibiting this property [AFGW10]. A good explanation of the generation process is provided in [BKMW10], which we summarize here. The generation process operates iteratively. It begins with an empty graph $G = (V, E) = (\emptyset, \emptyset)$ and progressively adds new vertices v_t to V , whose locations in the metric space are chosen randomly. Throughout this process, the generator maintains a series of 2^i -covers, denoted C_i , for each level i where $1 \leq i \leq \log D$, and D represents the diameter of the metric space. A set $C_i \subseteq V$ is a 2^i -cover if any two vertices $u, v \in C_i$ satisfy $d(u, v) \geq 2^i$, and every vertex $u \in V$ is within distance 2^i of some vertex in C_i . When a new vertex v_t is added, the generator identifies the smallest index i such that there exists a vertex $w \in C_i$ with $d(v_t, w) \leq 2^i$. The new vertex v_t is then added to all covers C_j for which $0 \leq j < i$. If no such



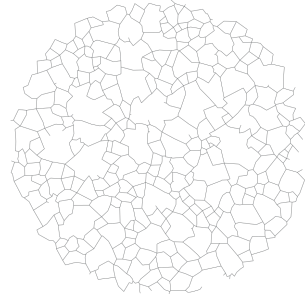
(a) Full Delaunay Triangulation (avg. deg. ≈ 6)



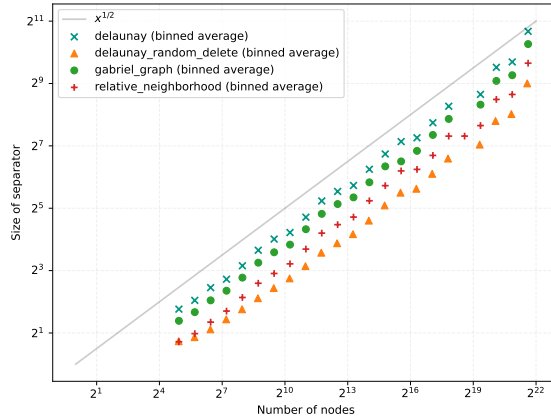
(b) Sparse DT (Random Deletion, avg. deg. ≈ 2.5)



(c) Gabriel Graph (GG)



(d) Relative Neighborhood Graph (RNG)



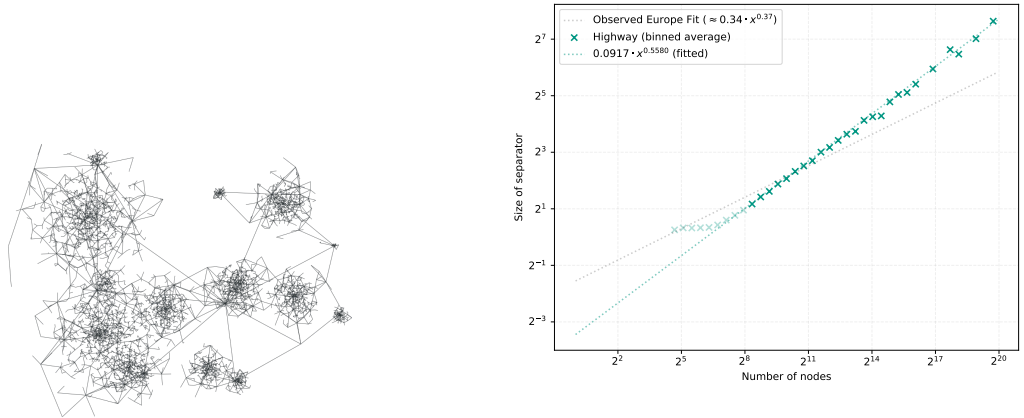
(e) Separator scaling for Delaunay variants.

Figure 5.9: Comparison of Delaunay Triangulation and its sparse subgraphs derived from the same point set: visualizations (a-d) and separator scaling (e).

index i exists, v_t is added to all cover sets C_j . Edges are subsequently added based on these covers and a tuning parameter k . For each cover C_j containing v_t (where $0 \leq j < i$), and for each existing vertex $w \in C_j$, an edge (w, v_t) is added if their distance satisfies $d(w, v_t) \leq k \cdot 2^j$. Furthermore, for each C_j containing v_t where $j < \log D$ and v_t is also present in C_{j+1} , an edge is added connecting v_t to its nearest neighbor within the set C_{j+1} . For our experiments, we adopt parameter settings similar to those used by [BKMW10], setting the diameter $D = 2^{25}$ and the connection parameter $k = \sqrt{2}$.

Bauer et al. also describe an alternative node sampling strategy aimed at creating structures resembling city clusters [BKMW10]. We implement and test both the uniform random sampling and this cluster-based sampling approach. Our experiments indicate no significant difference in the resulting separator sizes between the two sampling methods using the ABR generator framework.

The analysis of graphs generated using the ABR method yields separators that scale approximately as $\mathcal{O}(n^{1/2})$. This result is noteworthy because graphs generated by this process are typically highly non-planar. It serves as a reminder that observing $\mathcal{O}(n^{1/2})$ separator scaling does not necessarily imply planarity. Figure 5.10 provides a visual example of an ABR-generated graph and illustrates the observed separator scaling.



(a) Visualization of a synthetic graph generated using the ABR low highway dimension algorithm with 10^4 nodes.

(b) Separator size scaling observed during recursive partitioning of ABR-generated graphs. Separator sizes scale approximately as $\mathcal{O}(n^{1/2})$. The fit shown excludes subgraphs with ≤ 256 vertices due to their disproportionately large separators.

Figure 5.10: Analysis of synthetic graphs generated using the ABR algorithm [AFGW10].

5.5 Hierarchical Structure

In the following sections, we explore four distinct approaches to generating synthetic graphs with hierarchical structures.

5.5.1 Voronoi-Based Hierarchical Generator

One approach to generating synthetic networks with hierarchical features follows the method proposed by Bauer et al. [BKMW10]. This method utilizes Voronoi diagrams iteratively. A Voronoi diagram, given a set of points P , partitions the plane into convex regions, where each region contains the area closer to one point in P than to any other point in P . The generation process commences within a predefined initial polygon, which defines the spatial extent of the synthetic network. Points, designated as sites P , are then generated within this polygon according to a chosen sampling strategy. One strategy involves sampling a specified number of points uniformly at random throughout the polygon's interior. Alternatively, to better emulate settlement patterns, a clustered sampling approach can be employed. This approach involves first sampling locations for a number of population centers within the polygon. Each center is then assigned characteristics, such as a population size or an influence radius, drawn from a chosen distribution. Subsequently, points are sampled in the vicinity of each city center, concentrated within its influence radius, with the number of points related to the city's size.

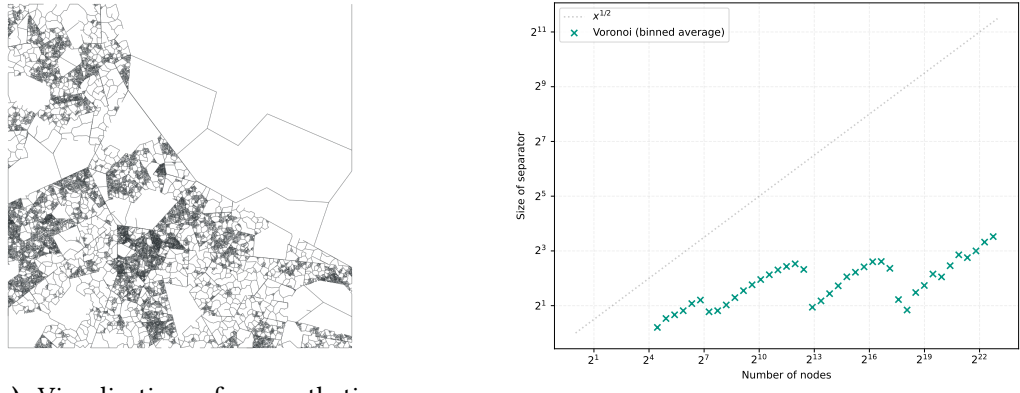
Regardless of the sampling method, the resulting set of points P serves as the input for computing the Voronoi diagram for this initial level. The Voronoi diagram partitions the polygon into regions based on nearest-neighbor relationships to the points in P . The edges of this Voronoi diagram (where adjacent regions meet) are then added to the synthetic graph. To introduce hierarchy, some of the resulting Voronoi regions (polygons) are selected, and the process is recursively applied within them: new points are sampled inside the selected region (using either uniform or clustered sampling), and a new Voronoi diagram is constructed within its boundaries, adding further edges to the graph.

Phase two of the generation process addresses the density of the graph G resulting from the Voronoi tessellation by constructing a sparse graph t -spanner H . A subgraph H is a t -spanner of G if it preserves all pairwise distances up to a multiplicative factor of t . The spanner is built using a greedy algorithm that iterates through the edges of G sorted by non-increasing length (longest first). An edge (u, v) with length $\text{len}(u, v)$ is added to the spanner H only if the current shortest path distance between u and v within H , denoted $\text{dist}_H(u, v)$, is greater than $t \cdot \text{len}(u, v)$. To compute $\text{dist}_H(u, v)$, Dijkstra's algorithm is employed. Additionally, a union-find data structure maintains the connected components of H , allowing for immediate addition of edges connecting previously disconnected vertices. Bauer et al. propose processing edges in non-increasing order (longest first) as a heuristic primarily intended to improve the computational efficiency of the spanner construction; the idea is to handle long edges while the intermediate spanner graph H is still sparse, potentially speeding up distance computations [BKMW10]. We observe that our pruning strategy, the specifics of which are presented in a later section in Algorithm 5.2, tends to yield graph structures more visually akin to real-world road networks compared to the Bauer et al. method, although this involves an increased computational expense.

This generation method introduces artifacts. For example, connections between major structures defined at higher levels (like large “cities”) might be unrealistically sparse, potentially consisting of only a few edges. Another artifact is that higher-level Voronoi edges (e.g., “highways” separating major regions) act as hard boundaries that lower-level edges (within regions) often do not cross, creating unrealistically effective separators along these top-level edges.

Analyzing separators in the final sparse graphs generated by this Voronoi-based method reveals interesting characteristics related to the hierarchy. Plots of separator size versus subgraph size often exhibit noticeable local minima, which appear to correspond to the

transitions between the hierarchical layers created during generation. This can lead to separators of approximately constant size when partitioning cuts primarily occur between these major structures at layer intersections. Within a single layer generated by the Voronoi tessellation (before sparsification), the separators tend to exhibit scaling closer to $\mathcal{O}(n^{1/2})$. Consequently, if the recursion depth is limited and the highest-level structures are allowed to grow large, their $\mathcal{O}(n^{1/2})$ separator behavior will dominate the separator size scaling. Figure 5.11 illustrates the structure and separator behavior of the final sparse graph.



(a) Visualization of a synthetic graph generated using the Voronoi method.

(b) Separator size scaling for the hierarchical Voronoi graph, showing local minima at layer transitions.

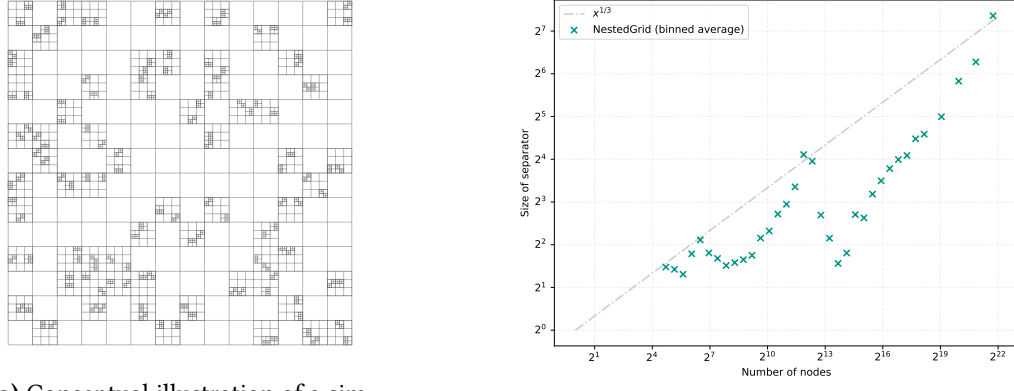
Figure 5.11: Analysis of synthetic graphs generated using the hierarchical Voronoi approach from Bauer et al. [BKMW10].

5.5.2 Nested Grids

To explore hierarchical effects in a more controlled manner, we also investigate a simpler but conceptually related approach using nested grids. This method aims to construct a hierarchical graph by recursively embedding grid graphs within the cells of a parent grid, potentially allowing us to deliberately engineer specific separator scaling properties. The fundamental idea involves placing copies of a subgraph (e.g., a square grid) into selected cells of a higher-level grid. This resulting composite graph can then itself be treated as a subgraph to be placed within cells of a grid at an even higher level, and so on. Figure 5.12 shows a small conceptual example of a nested grid with three layers.

A nested grid construction with l layers is determined by l grid sizes (specifying the dimensions of the grid at each layer) and $l - 1$ placement parameters (specifying how many subgraphs from layer $i+1$ are placed within layer i). This parameterization offers the possibility of attempting to enforce a target separator scaling, such as $\mathcal{O}(n^{1/3})$, by carefully choosing the parameters at each level transition. We assume that separating the nested structure primarily involves cutting through the top-level grid structure. If the number of embedded subgraphs is not excessively large, the separator size might be approximated by the width w of the top-level grid, similar to a standard grid separator. We can then estimate the total number of vertices $|V|$ at a given stage based on the grid width w of the current layer, the number k of subgraphs placed within its cells, and the size s of each subgraph:

$$|V| \approx w^2 + k \cdot s$$



(a) Conceptual illustration of a simple nested grid structure with three hierarchical levels.

(b) Separator size scaling for nested grids, showing pronounced local minima.

Figure 5.12: Analysis of synthetic graphs generated using the nested grid approach.

This estimate is approximate as it may double-count vertices along the boundaries where subgraphs connect to the parent grid. To enforce $\mathcal{O}(n^{1/3})$ scaling specifically at this layer transition, we can equate the separator size (approximated by w) with $|V|^{1/3}$:

$$w \approx (|V|)^{1/3} \approx (w^2 + ks)^{1/3}$$

This leads to the cubic equation $w^3 - w^2 - ks = 0$. If we fix the subgraph count k and subgraph size s , we can solve for the grid width w required to satisfy this condition at the transition. One root of this cubic equation for w is given by:

$$w = \frac{(27ks + \sqrt{(27ks + 2)^2 - 4 + 2})^{1/3}}{3 \cdot 2^{1/3}} + \frac{2^{1/3}}{3(27ks + \sqrt{(27ks + 2)^2 - 4 + 2})^{1/3}} + \frac{1}{3}$$

By carefully selecting w based on k and s using this relationship at each hierarchical level, one can attempt to construct a graph where the separator size scales roughly as the cube root of the number of nodes at each layer transition. This controlled approach allows the local minima in separator sizes, already seen in the Voronoi method, to be made more pronounced and regular, as all subgraphs at a given level transition can be designed with the same size s . However, achieving a consistent overall $\mathcal{O}(n^{1/3})$ scaling for the entire graph across all sizes n remains a challenge.

The observed local minima in separator size occur systematically at the transitions between hierarchical levels. This is because the construction method inherently creates sparse connections between components defined at different levels. Figure 5.13 provides a simplified illustration of this principle, showing two example components from the same levels within the hierarchy, not the entire structure of a layer. In this specific illustration, the components (represented by nodes 1-4 and 5-8) are linked only by the edges (3)-(5) and (4)-(6). Consequently, removing just one endpoint from each connecting edge suffices to disconnect them. This demonstrates how a vertex separator of small, constant size (size 2) exists between levels, regardless of the internal size of the components themselves. Such constant-size separators at every level transition are the underlying cause of the pronounced local minima observed in the separator scaling plots for these nested structures.

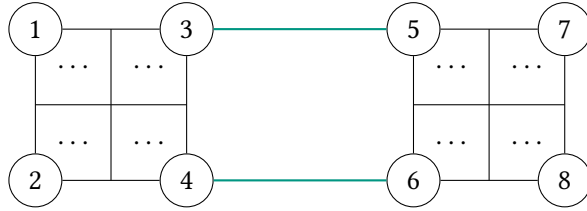


Figure 5.13: Simplified illustration of connectivity between components from adjacent hierarchical levels. The components (nodes 1–4 and 5–8) are connected only by the two green edges (3)–(5) and (4)–(6). Removing one endpoint from each edge (e.g., {3, 6}) forms a size-2 vertex separator, irrespective of component sizes.

5.5.3 Hierarchical Delaunay Graph Generation

Our efforts to generate synthetic graphs with realistic road network properties, particularly small separators, led us to refine techniques inspired by prior work on synthetic road network generation, such as that of Bauer et al. [BKMW10]. A significant artifact observed in some Voronoi-based hierarchical models was that major transport arteries, analogous to motorways, could not be realistically crossed by lower-level roads. This limitation arose because new expansion sites were generated strictly within the polygonal cells defined by the higher-level Voronoi tessellation, effectively making top-level Voronoi edges hard boundaries. Our revised core idea allows new centers of expansion to emerge from the existing network infrastructure itself; thus, new expansion sites are selected from the nodes of the current graph structure rather than from within predefined areal polygons.

An initial refined concept explored an iterative Delaunay approach. This process would commence with a global Delaunay triangulation of an initial point set. In subsequent hierarchical levels, a subset of existing points would be selected as *expansion sites*. New points would then be sampled in the vicinity of each of these sites. Following this, a new Delaunay triangulation would be performed, localized to the region around these newly added points and their respective expansion sites, and then integrated into the larger graph structure. This iterative process aimed to simulate a “rich get richer” phenomenon: regions with a higher density of points would be more likely to have their constituent points selected as expansion sites for subsequent levels, leading to further concentrated growth in those areas. This recursive generation would continue for a specified number of levels. As a final step, a planarization process, as described in Algorithm 4.1, would be applied: edges, interpreted as straight line segments, are checked for intersections. Any intersections found are resolved by introducing new vertices at these points and subdividing the original edges accordingly. This method showed initial promise in creating structures that could yield separators with scaling behavior similar to that of road networks.

Building upon insights from this iterative strategy, we developed a more streamlined and effective method, which is the primary focus of this section. Instead of performing Delaunay triangulations at each hierarchical level followed by a final planarization step, we first generate the complete set of points across all desired levels. Only after all points have been generated is a single, global Delaunay triangulation performed on the entire final point set. The pseudocode for this hierarchical Delaunay generation process is detailed in Algorithm 5.1.

A deliberate design choice in our hierarchical point generation process (Algorithm 5.1) is that candidates for new expansion sites at each level i are selected from the *entire* set of currently existing points P , rather than exclusively from points generated in the immediately

preceding level $i-1$. This approach reflects the real-world phenomenon where new settlements, even smaller ones, can emerge around and connect to established, major infrastructure hubs that might have formed at much earlier stages of network growth; for instance, a small town developing near a major motorway interchange. We also experimented with an alternative strategy where expansion sites for a given level were chosen only from the points generated in the previous level. However, this restriction did not yield any noticeable differences in the overall structural properties or separator characteristics of the resulting graphs, thus validating our choice of the more inclusive selection method.

Algorithm 5.1: Hierarchical Delaunay Graph Generation

Input: Number of hierarchical levels L ,
 Level expansion fractions f_i (where $f_1 = 1.0$),
 Points to generate per expansion site k_i ,
 Expansion radii r_i ,
 for $1 \leq i \leq L$

Output: Geometric graph G

```

1  $P \leftarrow \{p_{\text{random}}\}$ 
2 forall  $i \in [1, L]$  do
3    $C_i \leftarrow P.\text{chooseRandom}(\lfloor f_i \cdot |P| \rfloor)$ 
4   forall center  $\in C_i$  do
5      $P \leftarrow P \cup \text{randomPointsInCircle}(\text{center}, r_i, k_i)$ 
6  $G \leftarrow \text{delaunay}(P)$ 
7  $\text{pruneEdges}(G)$ 
8 return  $G$ 

```

Pruning Analogous to the delaunay triangulation approach in ??, the global Delaunay triangulation performed in Algorithm 5.1 results in a graph with an average vertex degree of approximately 6, which is denser than typical road networks.

To better emulate the characteristics of real road networks, such as an average degree around 2.5 and fewer nodes with very high degrees, we apply an edge pruning step. While Bauer et al. [BKMW10] also suggested a pruning methodology, we adopt a different strategy tailored to our generated graph structures. In our approach, edges are considered in order of increasing Euclidean length (shortest first). An edge (u, v) is removed if the shortest path distance between u and v in the graph, after the hypothetical removal of (u, v) , is no more than a factor α times the original direct length of (u, v) ; i.e., if $\text{dist}_{G \setminus \{(u, v)\}}(u, v) \leq \alpha \cdot \text{length}(u, v)$. This edge removal criterion approximates economic viability considerations, as connections that offer marginal utility or are largely redundant would likely not be constructed in real-world road networks. Through experimentation, we found that a pruning parameter of $\alpha = 2.5$ effectively reduces the average degree to the target range comparable to that of road networks. The pseudocode for this edge pruning process is given in Algorithm 5.2. A visual comparison of graph structures before and after pruning is provided in Figure 5.14.

The pruning step, while beneficial for achieving a target average degree and a more realistic degree distribution, does not fundamentally alter the asymptotic separator scaling of the generated Delaunay graphs. Figure 5.15 illustrates that the separator size scaling remains similar for both pruned and unpruned graphs.

Algorithm 5.2: Edge pruning based on path length redundancy.

Input: Graph $G = (V, E)$,
 Pruning parameter α
Output: Pruned version of G

```

1 forall  $(u, v) \in E$  sorted by increasing length do
2   if  $\text{dist}_{G \setminus \{(u, v)\}}(u, v) \leq \alpha \cdot \text{length}(u, v)$  then
3      $G \leftarrow G \setminus \{(u, v)\}$ 

```

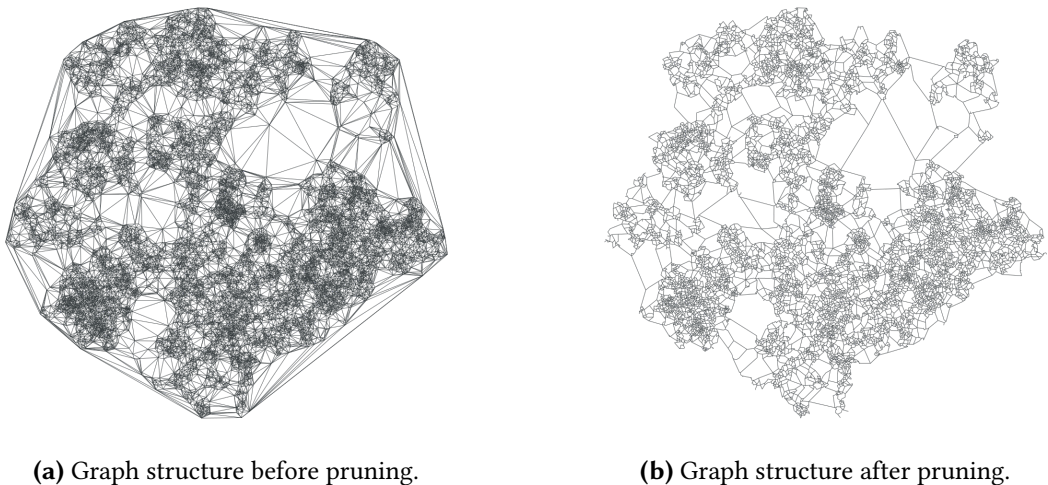


Figure 5.14: Visual comparison of a hierarchical Delaunay graph before and after edge pruning.

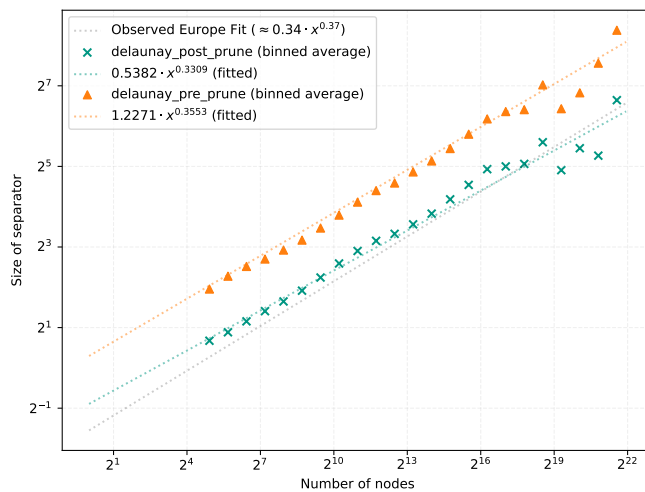


Figure 5.15: Comparison of separator size scaling in hierarchical Delaunay graphs before and after the pruning step.

Given that sequential pruning can be computationally intensive due to numerous shortest path computations (typically Dijkstra’s algorithm, where path precomputation is infeasible as the graph changes), we also explored an approximate parallel version. In this variant, edges are processed in chunks, often corresponding to the number of available processing threads. Shortest path computations for all candidate edges within a chunk are performed in parallel, based on the graph state *before* any edges in that chunk are removed. After all checks for the current chunk are complete, the identified edges are removed simultaneously. This parallelization introduces an approximation: an edge (u, v) might be removed because its shortest alternative path relied on another edge (x, y) from the same chunk, which is also concurrently identified for removal. If (x, y) is removed, the true shortest path for (u, v) in the updated graph might have become longer than the $\alpha \cdot \text{length}(u, v)$ threshold, but this would not be detected as the checks were based on the graph state at the beginning of the chunk processing. Despite this potential for over-pruning, our empirical results suggest that this approximation has a negligible impact on the final graph structure and its properties. This outcome is likely because edges are sorted by length before being chunked; thus, edges within a single chunk are often spatially distributed across the graph rather than being highly locally interdependent, minimizing negative interference from concurrent removals.

Parameterization The number of hierarchical layers, denoted L , is a key parameter for the generation process detailed in Algorithm 5.1. Drawing from observations of real road networks and prior work on synthetic graph generation [BKMW10], L is often set to 4, a choice suitable for generating graphs with up to approximately 20 million nodes. With $L = 4$, the algorithm requires parameters for level expansion fractions f_i , points per expansion site k_i , and expansion radii r_i , for $i \in [1, L]$, totaling $3L$ (i.e., 12 for $L = 4$) parameters. However, the effective number of free parameters is lower. The specific value of the first expansion radius, r_1 , primarily scales the entire embedding without fundamentally altering its topological structure, effectively reducing the count to 11 if r_1 is fixed or normalized. Furthermore, the algorithm specifies that the first level expansion fraction $f_1 = 1.0$ (ensuring the initial point always seeds the first level of expansion), reducing the free parameters to 10. If a target total number of vertices, n_{target} , is desired, one additional parameter (e.g., k_L) can be adjusted to meet this target, potentially leaving 9 free parameters for a 4-layer model. The total number of points after level j , denoted n_j (where $n_0 = 1$ is the initial seed point, so $n_1 = k_1 + 1$ represents the total points after the first expansion), is given by the recurrence relation:

$$n_j = \begin{cases} k_1 + 1 & \text{if } j = 1 \\ n_{j-1} + n_{j-1} \cdot f_j \cdot k_j & \text{if } j > 1 \text{ and } j \leq L \end{cases}$$

For example, for a 4-layer generation ($L = 4$), the total number of points n_4 is calculated iteratively:

$$\begin{aligned} n_1 &= k_1 + 1 \\ n_2 &= n_1 + n_1 \cdot f_2 \cdot k_2 \\ n_3 &= n_2 + n_2 \cdot f_3 \cdot k_3 \\ n_4 &= n_3 + n_3 \cdot f_4 \cdot k_4 \end{aligned}$$

The final graph size is thus n_L .

Parameter Interplay and Heuristics for Separator Properties Achieving road network-like separator scaling with the hierarchical Delaunay generator involves a nuanced interplay between the expansion fractions f_i , points per site k_i , and expansion radii r_i . A critical aspect for developing well-behaved separators appears to be ensuring sufficient interaction or *overlap* between regions expanded from different sites. This helps to avoid overly simplistic connections between hierarchical components, which can lead to pronounced local minima in separator scaling plots.

It is observed that decreasing expansion radii r_i contributes to desirable separator properties. The median edge length in a Delaunay triangulation of n points within a region of radius r scales as $r \cdot n^{-1/2}$. Thus, to maintain somewhat consistent local geometric scales as density increases within nested expansions (if k_i is relatively stable), r_i generally needs to decrease. Our empirical observations suggested that without an exponential decrease in radii across levels, achieving consistently small separators was not possible.

However, if radii become too small too quickly, the graph might develop insufficient connectivity between clusters originating from different parent sites. Consider a thought experiment: if each expansion site has an infinitesimally small influence radius for generating points for the next level, these new points primarily connect among themselves and back to their parent site from the higher level. The number of points on the effective “perimeter” of such an expanded site (those able to connect to other, distant parts of the graph) may grow much slower than the number of points generated internally, thereby keeping its external connectivity limited; this effect is amplified by the hierarchical sampling which tends to concentrate points internally. In such a scenario, one might find a separator for the graph defined by the top-level points (e.g., n_1 points after the first level) with a size scaling as $\mathcal{O}(n_1^{1/2})$. If each of these top-level separator nodes is a gateway to a large hierarchical substructure that is only sparsely connected back to this top-level framework (via few “perimeter” connections or the gateway node itself), the separator for the entire graph G_L might be estimated based on this top-level cut, extended by these few perimeter vertices. Given that the total number of nodes n_L is typically much larger than n_1 , such a separator (e.g., related to a $\mathcal{O}(n_1^{1/2})$ scaling) would likely be disproportionately small for n_L . This issue can also manifest at subsequent levels. An attempt to achieve a desired global $\mathcal{O}(n_L^{1/3})$ scaling can involve setting parameters so that the initial $\mathcal{O}(n_1^{1/2})$ separator scaling corresponds to this overall target. This, however, can lead to a different problem: distinct top-level expansion sites, even if spatially close, might end up connected by very few paths. This sparse inter-connectivity between major components results in the re-emergence of pronounced local minima in the separator scaling plots at transitions between hierarchical levels, an issue similar to that encountered in simpler nested grid or Voronoi generation schemes where components are too easily fractured. Therefore, for generating graphs with well-behaved separators, it is critical that the expansion areas originating from different sites *overlap* or interact sufficiently to build an integrated and robustly connected graph structure, rather than a collection of dense clusters weakly linked through a sparse higher-level skeleton.

The degree of spatial overlap between regions expanded from different sites is fundamental to achieving desirable separator characteristics, as it dictates the strength of connectivity between hierarchical components. The parameters f_i , k_i , and r_i jointly determine this overlap: A higher expansion fraction f_i results in a greater number of nodes from the parent level being selected as expansion sites. This reduces the average distance between chosen sites, causing their respective areas of influence for the current level to naturally adjoin or overlap significantly. Increasing k_i , the number of points generated per expansion site, primarily

possibly needs graphical illustration

makes each local cluster denser. Such high local density implies that partitioning these individual dense regions themselves requires larger separators, reflecting their increased internal complexity. Simultaneously, by populating the same expansion radius r_i with more points, the average distances between points decrease on this level, increasing the likelihood of inter-cluster connections. The effect of the expansion radius r_i is direct: a larger radius straightforwardly increases the spatial extent of each new cluster, leading to more substantial overlap with adjacent expansion zones and, consequently, also contributing to larger overall separators due to increased linkage. Radii at deeper hierarchical levels must also be scaled appropriately relative to those at higher levels to ensure desired structural properties and avoid unintended consequences on separator sizes. Therefore, a careful balance of these parameters, f_i , k_i , and r_i , is essential to cultivate a graph structure with the intended connectivity and separator properties.

The parameterization offers flexibility. For instance, to simulate prominent “urban centers” at higher levels, a larger initial radius r_1 might be employed. This would typically be balanced by adjusting subsequent parameters, such as reducing f_2 (selecting fewer centers from the points generated by these large initial expansions), to maintain overall structural integrity and desired separator characteristics. However, such compensations have practical limits; if radii do not shrink sufficiently across levels, adjustments to f_i or k_i may not fully mitigate adverse effects on separator scaling.

For generating structures visually akin to road networks, allowing the number of active expansion sites (primarily controlled by f_i) to decrease at deeper hierarchical levels often yields favorable results. This complements the inherent tendency of the generation process where all existing points are candidates for seeding new expansions (the “rich get richer” effect), helping to manage density while still fostering hierarchical differentiation. Plausible structures can emerge even with constant k_i values across levels, provided f_i and r_i are appropriately tuned. For instance, the parameter set comprising level expansion fractions $f_i = (1.0, 0.3, 0.2, 0.1)$, points per expansion site $k_i = (50, 50, 50, 50)$, and expansion radii $r_i = (1000, 424, 120, 17)$ produces graphs whose separator characteristics are illustrated in Figure 5.16.

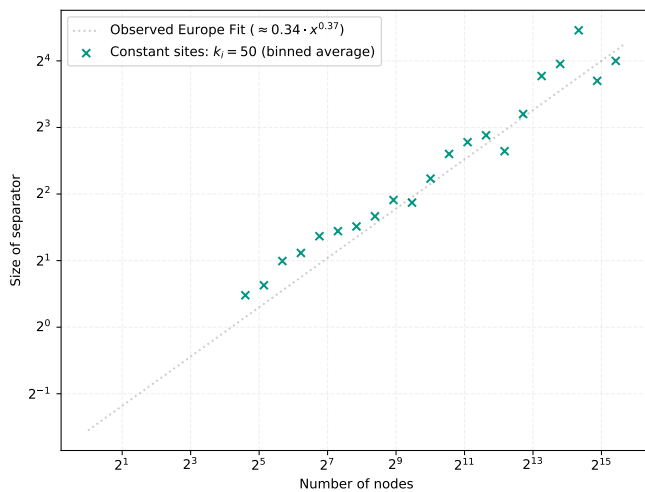


Figure 5.16: Separator size scaling for a hierarchical Delaunay graph generated with constant expansion sites: $f_i = (1.0, 0.3, 0.2, 0.1)$; $k_i = (50, 50, 50, 50)$; $r_i = (1000, 424, 120, 17)$.

To further illustrate the interplay of these parameters, Figure 5.17 presents separator scaling plots for four different parameter configurations. The baseline configuration, shown in Figure 5.17a, is chosen to produce separators with characteristics similar to those observed in real road networks. Figure 5.17b then demonstrates the impact of significantly decreasing the expansion fraction for the second level (f_2) while other parameters are held constant relative to the baseline; this results in a notable drop in separator sizes, indicative of a less interconnected structure. Subsequently, Figure 5.17c shows that this reduction in separator size due to a low f_2 can be counteracted by proportionally increasing the expansion radius of the second level (r_2) and the subsequent radii (r_3, r_4). Alternatively, Figure 5.17d illustrates that increasing the number of points generated at the first level (k_1) can also compensate for the reduced f_2 , again achieving separator scaling similar to the baseline by enhancing overall density and connectivity from the initial expansion phase.

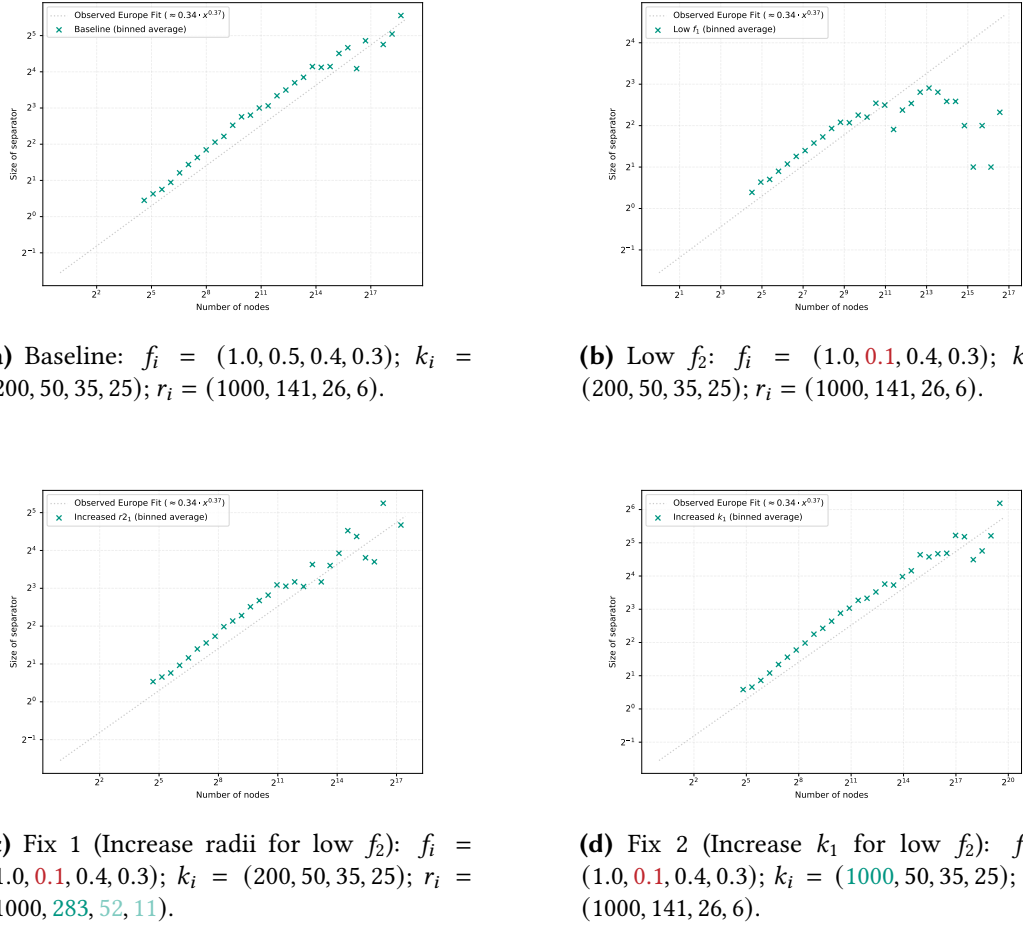


Figure 5.17: Illustration of parameter interplay on separator size scaling in hierarchical Delaunay graphs. Changed parameters in (b), (c), and (d) are relative to (a) or the specific modification being tested.

5.5.4 Physical Barriers

To investigate the influence of physical features of varying scales on road network structure and separator sizes, ranging from large-scale elements like mountains or lakes to smaller features like challenging terrain or rivers, we explore a generative approach based on iterating Perlin noise functions across multiple scales. The core idea is to use procedurally generated noise to define regions that are less favorable for node placement, simulating natural or man-made obstacles.

Our initial attempt utilizes a common method for generating additive fractal noise (often termed pink noise). This process involves summing multiple layers of Perlin noise, where at each step, the sampling frequency doubles and the amplitude halves. To avoid artifacts from sampling identical coordinates across different scales, we apply both scaling and translational offsets to the input coordinates for each noise layer: for instance, for a point (x, y) and a given scale, the noise is sampled at $(x \cdot \text{scale} + 3 \cdot \text{scale}, y \cdot \text{scale} + 3 \cdot \text{scale})$. If the scale values are chosen as powers of 2 adding ‘ $3 \cdot \text{scale}$ ’ to the coordinates ensures that the noise samples at different scales do not overlap. Points are then sampled in a domain, and their acceptance probability is modulated by the resulting smooth, cloud-like noise value at their location. However, this additive approach produces noise that is too smooth, and the resulting “obstacles” are not sufficiently “hard” or prohibitive. Graphs generated by accepting points based on this smooth additive fractal noise exhibit separator sizes scaling as $\mathcal{O}(n^{1/2})$.

To create more defined, “harder” obstacle boundaries across multiple scales, we develop an alternative approach using multiplicative Perlin noise. For each point p , Perlin noise is first sampled at M different scales (often referred to as octaves), where the frequency typically doubles at each successive scale. The output from the Perlin noise function for each of these M scales, usually in the range $[-1, 1]$, is then normalized to $[0, 1]$. Finally, the overall noise value for the point is determined by the product of these M normalized values. As more scales M are multiplied, this product naturally tends towards zero. To transform this continuous output into a binary map of allowed versus disallowed regions, we apply a noise threshold (e.g., 0.5^M): a point is considered in an “allowed” region if its final product value is greater than the noise threshold; otherwise, it is “disallowed”. This method yields a binary noise field with sharp boundaries and features at various frequencies, as Figure 5.18 illustrates.

One could also consider normalizing the product of the noise values by taking the M -th root of the product and using this value as a probability if a point is in an allowed region or not. This however yields quadratic scaling behavior for the separator size, as the resulting noise field is too smooth and does not produce sufficiently distinct obstacle regions.

The graph generation algorithm then utilizes this multiplicative binary Perlin noise to place vertices, as detailed in Algorithm 5.3. Essentially, candidate points are randomly sampled within a predefined domain (e.g., a unit circle). The noise value, which is the product of M scaled and normalized Perlin samples, is computed at each candidate point’s location. If this value indicates an “allowed” region, the point is accepted and added to the graph’s vertex set; otherwise, it is rejected, and a new candidate is sampled. This process continues until the desired number of n vertices is collected. For our experiments that yield good results for around 10 million nodes, we use a sequence of eight scales ($M = 8$): [8, 16, 32, 64, 128, 256, 512, 1024]. The initial scale of 8 is chosen instead of 1 because the Perlin noise implementation available to us exhibits overly smooth behavior at scale 1 for our purposes. Once the n points are generated, a global Delaunay triangulation is performed on this point set. Subsequently, the same edge pruning step detailed in the Hierarchical Delaunay Generation section (see

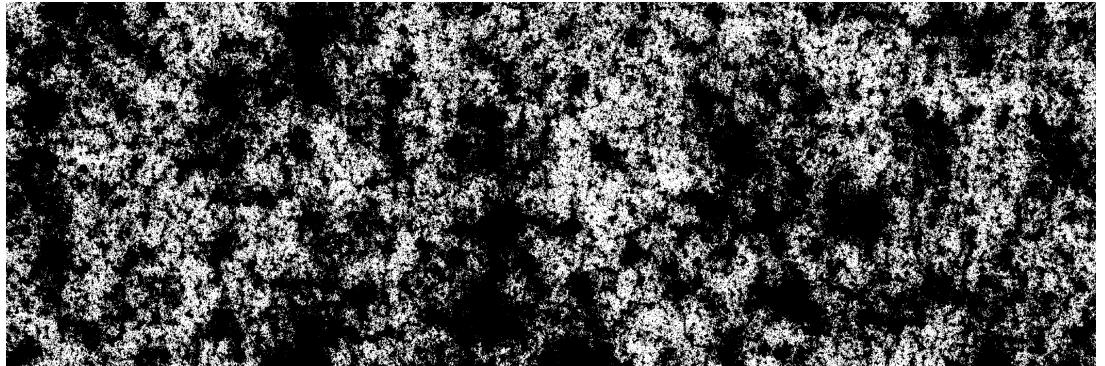


Figure 5.18: Visualization of the multiplicative binary Perlin noise used to define obstacle regions across multiple scales.

Algorithm 5.2) is applied to achieve a target average degree and refine the graph structure. An example of a graph generated using this method and its observed separator scaling are shown in Figure 5.19.

The specific sequence of noise scales is not universally optimal and should be adapted to the target graph size, n . For instance, using the aforementioned eight scales to generate a small graph of only 10^4 points results in separators that scale approximately as $\mathcal{O}(n^{1/2})$. This happens because, for a smaller total number of points, the high-frequency noise components can introduce excessive fine-grained fragmentation that obscures the structural influence of the larger, low-frequency obstacles. To achieve the desired separator behavior for smaller graphs, it is necessary to use fewer high-frequency scales, which allows the larger-scale features to more effectively shape the point distribution.

Algorithm 5.3: Graph generator using Multiplicative Binary Perlin Noise

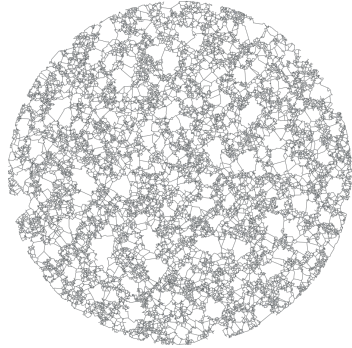
Input: Target number of points n ,
 M noise scales s_i
Output: Geometric graph G

```

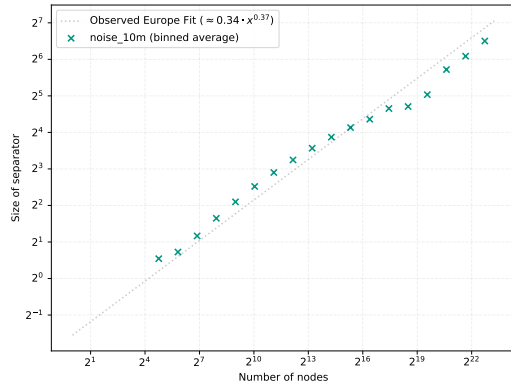
1  $P \leftarrow \emptyset$ 
2 while  $|P| < n$  do
3    $p \leftarrow \text{randomPointInUnitCircle}()$ 
4    $\text{noise} \leftarrow \prod_{i=1}^M \text{perlinNoise}(p \cdot s_i)$ 
5   if  $\text{noise} > 0.5^M$  then
6      $P \leftarrow P \cup \{p\}$ 
7    $G \leftarrow \text{delaunay}(P)$ 
8    $\text{pruneEdges}(G)$ 
9 return  $G$ 

```

This generation process does not directly prevent Delaunay edges from spanning the “obstacle” regions (areas with low composite noise values where no points are placed). However, it ensures that no vertices, and thus no potential navigation hubs or major junctions, are located within these simulated forbidden zones. Furthermore, the subsequent edge pruning



(a) Graph generated using multi-scale Perlin noise-based point sampling ($n = 10^5$).



(b) Separator scaling for the multi-scale Perlin noise graph.

Figure 5.19: Synthetic graph generated using multiplicative binary Perlin noise and its separator scaling.

step tends to remove many of the long, redundant edges that might span these empty regions, preserving only those deemed more critical for connectivity, akin to bridges over rivers or passes through mountainous terrain.

An interesting finding is that this multi-scale Perlin noise generation method, requiring only selection of a suitable range of scales rather than extensive parameter fine-tuning, produces graphs whose separator sizes scale approximately as $\mathcal{O}(n^{0.37})$. This is particularly noteworthy as an exponent of approximately 0.37 aligns closely with the separator scaling behavior empirically observed in real-world road networks, being less than $\mathcal{O}(n^{1/3})$, yet also substantially better than $\mathcal{O}(n^{1/2})$.

Ablation Studies on Noise Scales We also investigate whether the full spectrum of noise scales is necessary to achieve this result. Experiments are conducted using only high-frequency scales ([128, 256, 512, 1024]) and only low-frequency scales ([8, 16, 32, 64]). The resulting noise patterns and separator scaling comparisons are illustrated in Figure 5.20. When using only high-frequency scales, the separators for smaller subgraphs (identified through nested dissection) align well with the desired trend. However, as the graph size or region under consideration increases, the separator sizes revert to an $\mathcal{O}(n^{1/2})$ trend, because large-scale obstacle features (which would be defined by low frequencies) are absent. Conversely, when using only low-frequency scales, the separator behavior changes with the graph’s size. For smaller subgraphs, which mostly fit within the open areas created by high noise values, separators tend to scale as $\mathcal{O}(n^{1/2})$; this is similar to what is expected from standard Delaunay triangulations in unobstructed regions. However, as subgraphs become larger and the low-frequency obstacles themselves begin to define the separations, a “dropoff” from this initial $\mathcal{O}(n^{1/2})$ trend is observed, meaning the separators become relatively smaller for these intermediate sizes.

These studies therefore show that using a combination of multiple noise scales, representing obstacles of various sizes, is key to consistently achieving small separator sizes across a wide range of graph dimensions with this multi-scale Perlin noise method. This reliance on features defined across different scales to create a complex, layered obstacle landscape is

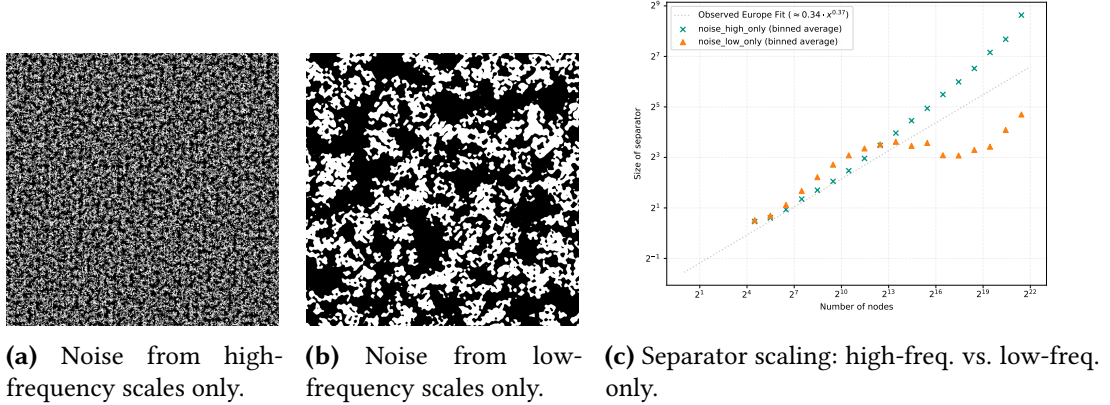


Figure 5.20: Impact of using only high-frequency versus only low-frequency Perlin noise scales on (a, b) noise patterns and (c) separator scaling.

conceptually similar to the explicit hierarchical graph generation methods explored earlier (e.g., Section 5.5.3); in both approaches, such multi-scale or hierarchical structuring is vital for achieving the desired separator properties, as is demonstrated in our noise model experiments where the favorable separator scaling breaks down if an insufficient range of scales is employed.

Alternative Graph Construction: k-Nearest Neighbors While the primary approach described above utilizes a Delaunay triangulation followed by pruning on the noise-sampled points, we also explored an alternative graph construction method using k-Nearest Neighbors (k-NN) on the same underlying point sets. For these non-uniformly distributed points, a consequence of the noise-based acceptance criteria, a $k = 3$, which often suffices to ensure reasonable connectivity for uniformly sampled points, proved insufficient. We found that a value of $k = 5$ was generally necessary to achieve a largely connected graph. Interestingly, k-NN graphs constructed from these noise-sampled points exhibited separator sizes that were substantially smaller than our target $\mathcal{O}(n^{0.37})$. Figure 5.21 illustrates this comparison. Visually comparing a k-NN graph (Figure 5.21b) with a graph derived from the Delaunay-based approach on the same noise-sampled point set (Figure 5.21a) reveals that the k-NN graph possesses significantly fewer long edges. This characteristic arises because the k-NN construction inherently limits connections to only the k closest neighbors. Consequently, edges are less likely to span large low-noise regions (our simulated obstacles), particularly if a point's k nearest neighbors all reside on the same side of such a zone. Marginally increasing k does not fundamentally alter this effect, as it still restricts connections to a local neighborhood. The limited long-range connectivity of the k-NN rule appears to cascade through recursive partitioning. This leads to disproportionately small separators for larger graphs, while smaller, more internally-connected subgraphs can exhibit comparatively larger separators. For substantially larger values of k , such as ($k=50$), the separators exhibit quadratic scaling.

write something like: we need hierarchy + delaunay that can span larger distances to get the desired scaling

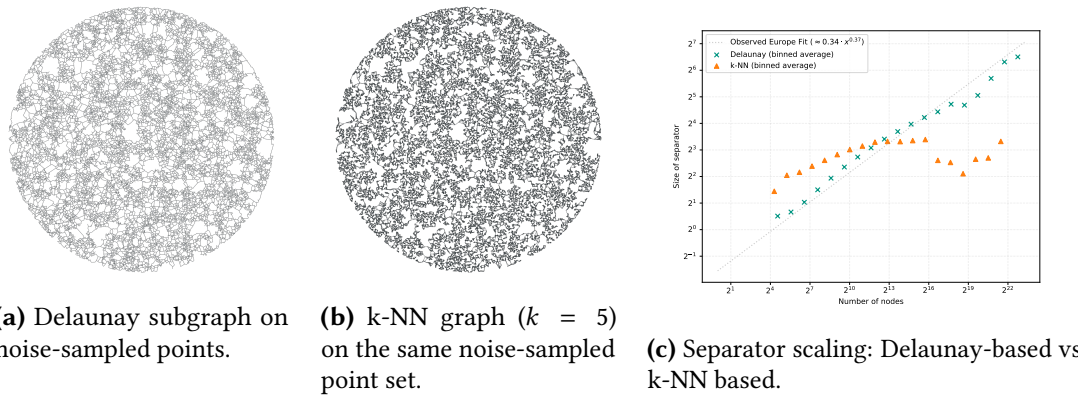


Figure 5.21: Comparison of graph structures and separator scaling for Delaunay-derived graphs versus k-NN graphs on identical noise-sampled point sets.

6 Conclusion

This thesis investigates the underlying structural properties responsible for the small separators observed in road networks.

6.1 Summary of Findings

Our empirical analysis first refines the understanding of this phenomenon. We demonstrate that the separator size s scales with graph size n according to the power law $s \approx \mathcal{O}(n^{0.37})$. This exponent is slightly larger than the previously suggested $\mathcal{O}(n^{1/3})$ [DSW16] but remains substantially better than the $\mathcal{O}(n^{1/2})$ bound for planar graphs.

Our investigation systematically demonstrates the insufficiency of simple, isolated graph properties to explain this behavior. Models based solely on degree distribution or basic geometric locality consistently produce separators that scale as $\mathcal{O}(n)$ or $\mathcal{O}(n^{1/2})$. Even when considering planarity, a defining feature of road networks, standard models like sparse grids or simple Delaunay triangulations fail to reproduce the target scaling. The planarization of real road networks, furthermore, has a negligible impact on their separator sizes, suggesting that the non planarity nature of these graphs is not a primary factor in the observed small separators.

In contrast, models incorporating structure across multiple scales show significant promise. Our proposed Hierarchical Delaunay generator constitutes an entire parameterized class of graphs. Through careful tuning of its parameters, governing expansion fractions, points per site, and radii across multiple levels, this model is also capable of replicating the observed $\mathcal{O}(n^{0.37})$ scaling. Notably, the resulting graph structures often bear a strong visual resemblance to those generated by the physical barrier model.

The most successful synthetic model developed in this thesis, however, is one based on simulating physical barriers using multi-scale Perlin noise. This approach generates a complex, layered landscape of obstacles at various scales, constraining where vertices can be placed. Remarkably, graphs generated using this method, which combines noise-based point sampling with a final Delaunay triangulation and pruning, naturally produce separators that scale approximately as $\mathcal{O}(n^{0.37})$. This result is achieved without extensive parameter fine-tuning, suggesting that this model captures a more fundamental generative principle. Ablation studies further underscore this finding, demonstrating that the full spectrum of noise scales, from large, regional barriers to small, local ones, is crucial for achieving this specific scaling across a wide range of graph sizes.

It is important to qualify, however, that the need for a multi-scale or hierarchical structure is not an absolute constraint if one allows for extreme parameter fine-tuning. The tree-locality model, for instance, could replicate the desired scaling using a highly specific distance-decay function $f(\text{dist}) = 1/\text{dist}^{3.3}$. This case serves as an interesting exception, highlighting that specific correlation structures can be enforced without explicit hierarchy, though this offers less insight into the natural emergence of such properties.

Revisiting the central research question “Do small separators in road networks arise from intrinsic graph properties, or from real-world physical barriers?” our findings provide a nuanced answer. The evidence strongly suggests that small separators are not a consequence of any single, simple graph property but are an emergent feature of a complex, multi-scale structural organization. This multi-scale structure can be interpreted as an explicit *hierarchy* of road types or, perhaps more fundamentally, as the result of *physical barriers* existing at all geographic scales. The success of the multi-scale Perlin noise model indicates that the constrained placement of nodes and edges by geography is a powerful generative mechanism. Road networks are built upon a landscape permeated by obstacles at every scale, from mountain ranges down to local properties. This forces the graph to be composed of densely connected regions that are themselves sparsely interconnected through well-defined corridors, a structure highly amenable to small separators. We therefore conclude that the small separators in road networks are a direct result of their adaptation to a multi-scale, obstacle-rich environment.

6.2 Limitations and Future Work

The conclusions presented in this thesis are primarily derived from experimental analysis. While our results strongly indicate the importance of hierarchy and multi-scale barriers, the inability of simpler, non-hierarchical models to reproduce the desired scaling does not constitute a formal disproof of their potential under different assumptions. A notable limitation is that our most successful models rely on a final Delaunay triangulation. The observation that k-Nearest Neighbor graphs built on the same hierarchical point sets fail to produce the correct scaling suggests that the specific connectivity pattern of Delaunay graphs, which allows for both local and some structured long-range connections, plays a significant, yet not fully explored, role.

These limitations and findings open several avenues for future research. A primary direction is the pursuit of a rigorous theoretical analysis of the multiplicative Perlin noise model. Such a study could potentially provide a formal derivation for the empirically observed $\mathcal{O}(n^{0.37})$ scaling. Further work could also seek to improve the efficiency of the generative algorithms. For instance, the tree-locality generator is hampered by its slow, BFS-based distance calculations; developing an efficient method to sample edges according to weights derived from tree distances would make this model more viable for large-scale tests. Finally, the synthetic generators developed here can serve as a basis for creating realistic, large-scale benchmarks for evaluating not only route planning algorithms but also other algorithms that depend on network topology. Exploring alternative models for generating hierarchical structures or physical barriers could yield further insights into the essential characteristics of transportation networks.

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