ONCE UPON A TIME PEOPLE WERE BORN INTO COMMUNITIES AND HAD TO FIND THEIR INDIVIDUALITY. TODAY PEOPLE ARE BORN INDIVIDUALS AND HAVE TO FIND THEIR COMMUNITIES.

MASS INDIE RESPONDS TO THIS SITUATION BY CREATING CLIQUES OF PEOPLE IN THE KNOW, WHILE NORMCORE KNOWS THE REAL FEAT IS HARRNESSING THE POTENTIAL FOR CONNECTION TO SPING UP. IT’S ABOUT ADAPABILITY, NOT EXCLUSIVITY.
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**Figure 9.0 (cover image)**

**Art & Networks: K-Mode: Youth Mode**

K-Mode is an art collective that publishes trend reports with an unusual take on various concepts. The image shows a page from *Youth Mode: A Report on Freedom*, discussing the subtle shift in the origins and the meaning of communities, the topic of this chapter [1].
INTRODUCTION

Belgium appears to be the model bicultural society: 59% of its citizens are Flemish, speaking Dutch and 40% are Walloons who speak French. As multiethnic countries break up all over the world, we must ask: How did this country foster the peaceful coexistence of these two ethnic groups since 1830? Is Belgium a densely knitted society, where it does not matter if one is Flemish or Walloon? Or we have two nations within the same borders, that learned to minimize contact with each other?

The answer was provided by Vincent Blondel and his students in 2007, who developed an algorithm to identify the country’s community structure. They started from the mobile call network, placing individuals next to whom they regularly called on their mobile phone [2]. The algorithm revealed that Belgium’s social network is broken into two large clusters of communities and that individuals in one of these clusters rarely talk with individuals from the other cluster (Figure 9.1). The origin of this separation became obvious once they assigned to each node the language spoken by each individual, learning that one cluster consisted almost exclusively of French speakers and the other collected the Dutch speakers.

In network science we call a community a group of nodes that have a higher likelihood of connecting to each other than to nodes from other communities. To gain intuition about community organization, next we discuss two areas where communities play a particularly important role:

- **Social Networks**
  Social networks are full of easy to spot communities, something that scholars have noticed decades ago [3,4,5,6,7]. Indeed, the employees of a company are more likely to interact with their coworkers than with employees of other companies [3]. Consequently work places appear as densely interconnected communities within the social network. Communities could also represent circles of friends, or a group of individuals who pursue the same hobby together, or individuals living in the same neighborhood.
  A social network that has received particular attention in the context...
The interest in the dataset is driven by a singular event: A conflict between the club’s president and the instructor split the club into two. About half of the members followed the instructor and the other half the president, a breakup that unveiled the ground truth, representing club’s underlying community structure (Figure 9.2a). Today community finding algorithms are often tested based on their ability to infer these two communities from the structure of the network before the split.

**Biological Networks**

Communities play a particularly important role in our understanding of how specific biological functions are encoded in cellular networks. Two years before receiving the Nobel Prize in Medicine, Lee Hartwell argued that biology must move beyond its focus on single genes. It must explore instead how groups of molecules form functional modules to carry out a specific cellular functions [10]. Ravasz and collaborators [11] made the first attempt to systematically identify such modules in metabolic networks. They did so by building an algorithm to identify groups of molecules that form locally dense communities (Figure 9.3).

Communities play a particularly important role in understanding human diseases. Indeed, proteins that are involved in the same disease tend to interact with each other [12,13]. This finding inspired the disease module hypothesis [14], stating that each disease can be linked to a well-defined neighborhood of the cellular network.

The examples discussed above illustrate the diverse motivations that drive community identification. The existence of communities is rooted in who connects to whom, hence they cannot be explained based on the degree distribution alone. To extract communities we must therefore inspect a network’s detailed wiring diagram. These examples inspire the starting hypothesis of this chapter:

H1: Fundamental Hypothesis

A network’s community structure is uniquely encoded in its wiring diagram.

According to the fundamental hypothesis there is a ground truth about a network’s community organization, that can be uncovered by inspecting $A_{ij}$.

The purpose of this chapter is to introduce the concepts necessary to...
understand and identify the community structure of a complex network. We will ask how to define communities, explore the various community characteristics and introduce a series of algorithms, relying on different principles, for community identification.

![Image](image-url)

**Figure 9.3**

**Communities in Metabolic Networks**

The E. coli metabolism offers a fertile ground to investigate the community structure of biological systems [11].

(a) The biological modules (communities) identified by the Ravasz algorithm [11] (SECTION 9.3). The color of each node, capturing the predominant biochemical class to which it belongs, indicates that different functional classes are segregated in distinct network neighborhoods. The highlighted region selects the nodes that belong to the pyrimidine metabolism, one of the predicted communities.

(b) The topologic overlap matrix of the E. coli metabolism and the corresponding dendrogram that allows us to identify the modules shown in (a). The color of the branches reflect the predominant biochemical role of the participating molecules, like carbohydrates (blue), nucleotide and nucleic acid metabolism (red), and lipid metabolism (cyan).

(c) The red right branch of the dendrogram tree shown in (b), highlighting the region corresponding to the pyridine module.

(d) The detailed metabolic reactions within the pyrimidine module. The boxes around the reactions highlight the communities predicted by the Ravasz algorithm.

What do we really mean by a community? How many communities are in a network? How many different ways can we partition a network into communities? In this section we address these frequently emerging questions in community identification.

DEFINING COMMUNITIES
Our sense of communities rests on a second hypothesis (Figure 9.4):

**H2: Connectedness and Density Hypothesis**

A community is a locally dense connected subgraph in a network.

In other words, all members of a community must be reached through other members of the same community (connectedness). At the same time we expect that nodes that belong to a community have a higher probability to link to the other members of that community than to nodes that do not belong to the same community (density). While this hypothesis considerably narrows what would be considered a community, it does not uniquely define it. Indeed, as we discuss below, several community definitions are consistent with H2.

**Maximum Cliques**

One of the first papers on community structure, published in 1949, defined a community as group of individuals whose members all know each other [5]. In graph theoretic terms this means that a community is a complete subgraph, or a clique. A clique automatically satisfies H2: it is a connected subgraph with maximal link density. Yet, viewing communities as cliques has several drawbacks:

- While triangles are frequent in networks, larger cliques are rare.
- Requiring a community to be a complete subgraph may be too restrictive, missing many other legitimate communities. For example, none of the communities of Figure 9.2 and 9.3 correspond to complete subgraphs.
**Strong and Weak Communities**

To relax the rigidity of cliques, consider a connected subgraph \( C \) of \( N_c \) nodes in a network. The **internal degree** \( k_{i\text{int}} \) of node \( i \) is the number of links that connect \( i \) to other nodes in \( C \). The **external degree** \( k_{i\text{ext}} \) is the number of links that connect \( i \) to the rest of the network. If \( k_{i\text{ext}} = 0 \), each neighbor of \( i \) is within \( C \), hence \( C \) is a good community for node \( i \). If \( k_{i\text{int}} = 0 \), then node \( i \) should be assigned to a different community. These definitions allow us to distinguish two kinds of communities (Figure 9.5):

- **Strong Community**
  
  \( C \) is a strong community if each node within \( C \) has more links within the community than with the rest of the graph \([15,16]\). Specifically, a subgraph \( C \) forms a strong community if for each node \( i \in C \),
  
  \[
  k_{i\text{int}}(C) > k_{i\text{ext}}(C).
  \]  

  (9.1)

- **Weak Community**
  
  \( C \) is a weak community if the total internal degree of a subgraph exceeds its total external degree \([16]\). Specifically, a subgraph \( C \) forms a weak community if
  
  \[
  \sum_{i \in C} k_{i\text{int}}(C) > \sum_{i \in C} k_{i\text{ext}}(C).
  \]  

  (9.2)

A weak community relaxes the strong community requirement by allowing some nodes to violate (9.1). In other words, the inequality (9.2) applies to the community as a whole rather than to each node individually.

Note that each clique is a strong community, and each strong community is a weak community. The converse is generally not true (Figure 9.5).

The community definitions discussed above (cliques, strong and weak communities) refine our notions of communities. At the same time they indicate that we do have some freedom in defining communities.

**NUMBER OF COMMUNITIES**

How many ways can we group the nodes of a network into communities? To answer this question consider the simplest community finding problem, called **graph bisection**: We aim to divide a network into two non-overlapping subgraphs, such that the number of links between the nodes in the two groups, called the cut size, is minimized (BOX 9.1).

**Graph Partitioning**

We can solve the graph bisection problem by inspecting all possible divisions into two groups and choosing the one with the smallest cut size. To determine the computational cost of this brute force approach we note that the number of distinct ways we can partition a network of \( N \) nodes into groups of \( N_1 \) and \( N_2 \) nodes is

\[
\frac{N!}{N_1!N_2!}.
\]  

(9.3)
Chip designers face a problem of exceptional complexity: They need to place on a chip 2.5 billion transistors such that their wires do not intersect. To simplify the problem they first partition the wiring diagram of an integrated circuit (IC) into smaller subgraphs, chosen such that the number of links between them to be minimal. Then they lay out different blocks of an IC individually, and reconnect these blocks. A similar problem is encountered in parallel computing, when a large computational problem is partitioned into subtasks and assigned to individual chips. The assignment must minimize the typically slow communication between the processors.

The problem faced by chip designers or software engineers is called graph partitioning in computer science [17]. The algorithms developed for this purpose, like the widely used Kerninghan-Lin algorithm (Figure 9.6), are the predecessors of the community finding algorithms discussed in this chapter.

There is an important difference between graph partitioning and community detection: Graph partitioning divides a network into a predefined number of smaller subgraphs. In contrast community detection aims to uncover the inherent community structure of a network. Consequently in most community detection algorithms the number and the size of the communities is not predefined, but needs to be discovered by inspecting the network’s wiring diagram.

Using Stirling’s formula \( n! = \sqrt{2\pi n}(n/e)^n \) we can write (9.3) as

\[
\frac{N!}{N_1!N_2!} \approx \frac{\sqrt{2\pi N}(N/e)^N}{\sqrt{2\pi N_1}(N_1/e)^{N_1}} \frac{\sqrt{2\pi N}(N/e)^N}{\sqrt{2\pi N_2}(N_2/e)^{N_2}} \approx \frac{N^{N+1/2}}{N_1^{N_1+1/2}N_2^{N_2+1/2}}. \tag{9.4}
\]

To simplify the problem let us set the goal of dividing the network into two equal sizes \( N_1 = N_2 = N/2 \). In this case (9.4) becomes

\[
\frac{2^{N+1}}{\sqrt{N}} \approx e^{N+1/4} - \frac{1}{2N}, \tag{9.5}
\]

indicating that the number of bisections increases exponentially with the size of the network.

To illustrate the implications of (9.5) consider a network with ten nodes
which we bisect into two subgraphs of size \( N_1 = N_2 = 5 \). According to (9.3) we need to check 252 bisections to find the one with the smallest cut size. Let us assume that our computer can inspect these 252 bisections in one millisecond (\( 10^{-3} \) sec). If we next wish to bisect a network with a hundred nodes into groups with \( N_1 = N_2 = 50 \), according to (9.3) we need to check approximately \( 10^{29} \) divisions, requiring about \( 10^{16} \) years on the same computer. Therefore our brute-force strategy is bound to fail, being impossible to inspect all bisections for even a modest size network.

**Community Detection**

While in graph partitioning the number and the size of communities is predefined, in community detection both parameters are unknown. We call a partition a division of a network into an arbitrary number of groups, such that each node belongs to one and only one group. The number of possible partitions follows [19-22]

\[
B_N = \frac{1}{e} \sum_{j=0}^{\infty} \frac{j^N}{j!}.
\]

As Figure 9.7 indicates, \( B_N \) grows faster than exponentially with the network size for large \( N \).

Equations (9.5) and (9.6) signal the fundamental challenge of community identification: The number of possible ways we can partition a network into communities grows exponentially or faster with the network size \( N \). Therefore it is impossible to inspect all partitions of a large network (BOX 9.2).

In summary, our notion of communities rests on the expectation that each community corresponds to a locally dense connected subgraph. This hypothesis leaves room for numerous community definitions, from cliques to weak and strong communities. Once we adopt a definition, we could identify communities by inspecting all possible partitions of a network, selecting the one that best satisfies our definition. Yet, the number of partitions grows faster than exponentially with the network size, making such brute-force approaches computationally infeasible. We therefore need algorithms that can identify communities without inspecting all partitions. This is the subject of the next sections.
NP COMPLETENESS

How long does it take to execute an algorithm? The answer is not given in minutes and hours, as the execution time depends on the speed of the computer on which we run the algorithm. We count instead the number of computations the algorithm performs. For example an algorithm that aims to find the largest number in a list of $N$ numbers has to compare each number in the list with the maximum found so far. Consequently its execution time is proportional to $N$. In general, we call an algorithm polynomial if its execution time follows $N^p$.

An algorithm whose execution time is proportional to $N^3$ is slower on any computer than an algorithm whose execution time is $N$. But this difference dwindles in significance compared to an exponential algorithm, whose execution time increases as $2^N$. For example, if an algorithm whose execution time is proportional to $N$ takes a second for $N = 100$ elements, then an $N^3$ algorithm takes almost three hours on the same computer. Yet an exponential algorithm ($2^N$) will take $10^{20}$ years to complete.

The problem that an algorithm can solve in polynomial time is called a class $P$ problem. Several computational problems encountered in network science have no known polynomial time algorithms, but the available algorithms require exponential running time. Yet, the correctness of the solution can be checked quickly, i.e. in polynomial time. Such problems, called NP-complete, include the traveling salesman problem (Figure 9.8), the graph coloring problem, maximum clique identification, partitioning a graph into subgraphs of specific type, and the vertex cover problem (Box 7.4).

The ramifications of NP-completeness has captured the fascination of the popular media as well. Charlie Epps, the main character of the CBS TV series Numbers, spends the last three months of his mother’s life trying to solve an NP complete problem, convinced that the solution will cure her disease. Similarly the motive for a double homicide in the CBS TV series Elementary is the search for a solution of an NP-complete problem, driven by its enormous value for cryptography.
To uncover the community structure of large real networks we need algorithms whose running time grows polynomially with $N$. Hierarchical clustering, the topic of this section, helps us achieve this goal.

The starting point of hierarchical clustering is a similarity matrix, whose elements $x_{ij}$ indicate the distance of node $i$ from node $j$. In community identification the similarity is extracted from the relative position of nodes $i$ and $j$ within the network.

Once we have $x_{ij}$, hierarchical clustering iteratively identifies groups of nodes with high similarity. We can use two different procedures to achieve this: agglomerative algorithms merge nodes with high similarity into the same community, while divisive algorithms isolate communities by removing low similarity links that tend to connect communities. Both procedures generate a hierarchical tree, called a dendrogram, that predicts the possible community partitions. Next we explore the use of agglomerative and divisive algorithms to identify communities in networks.

AGGLOMERATIVE PROCEDURES: THE RAVASZ ALGORITHM

We illustrate the use of agglomerative hierarchical clustering for community detection by discussing the Ravasz algorithm, proposed to identify functional modules in metabolic networks [11]. The algorithm consists of the following steps:

Step 1: Define the Similarity Matrix

In an agglomerative algorithm similarity should be high for node pairs that belong to the same community and low for node pairs that belong to different communities. In a network context nodes that connect to each other and share neighbors likely belong to the same community, hence their $x_{ij}$ should be large. The topological overlap matrix (Figure 9.9) [11]

$$x_{ij}^0 = \frac{J(i, j)}{\min(k_i, k_j) + 1 - \Theta(A_{ij})}$$

(9.7)
captures this expectation. Here $\Theta(x)$ is the Heaviside step function, which is zero for $x \leq 0$ and one for $x > 0$; $l(i,j)$ is the number of common neighbors of node $i$ and $j$, to which we add one (+1) if there is a direct link between $i$ and $j$; $\min(k_i,k_j)$ is the smaller of the degrees $k_i$ and $k_j$. Consequently:

- $x_{ij}^0 = 1$ if nodes $i$ and $j$ have a link to each other and have the same neighbors, like A and B in Figure 9.9a.
- $x_{ij}^0 = 0$ if $i$ and $j$ do not have common neighbors, nor do they link to each other, like A and E.
- Members of the same dense local network neighborhood have high topological overlap, like nodes H, I, J, K or E, F, G.

**Step 2: Decide Group Similarity**

As nodes are merged into small communities, we must measure how similar two communities are. Three approaches, called single, complete and average cluster similarity, are frequently used to calculate the community similarity from the node-similarity matrix $x_{ij}$ (Figure 9.10). The Ravasz algorithm uses the average cluster similarity method, defining the similarity of two communities as the average of $x_{ij}$ over all node pairs $i$ and $j$ that belong to distinct communities (Figure 9.10d).

**Step 3: Apply Hierarchical Clustering**

The Ravasz algorithm uses the following procedure to identify the communities:

1. Assign each node to a community of its own and evaluate $x_{ij}$ for all node pairs.
2. Find the community pair or the node pair with the highest similarity and merge them into a single community.
3. Calculate the similarity between the new community and all other communities.
4. Repeat Steps 2 and 3 until all nodes form a single community.

**Step 4: Dendrogram**

The agglomerative hierarchical clustering algorithm proposed by Ravasz was designed to identify functional modules in metabolic networks, but it can be applied to arbitrary networks.

(a) **Topological Overlap**

A small network illustrating the calculation of the topological overlap $x_{ij}^0$. For each node pair $i$ and $j$ we calculate the overlap $x_{ij}^0$. The obtained $x_{ij}^0$ for each connected node pair is shown on each link. Note that $x_{ij}^0$ can be nonzero for nodes that do not link to each other, but have a common neighbor. For example, $x_{ij}^0 = 1/3$ for C and E.

(b) **Topological Overlap Matrix**

The topological overlap matrix $x_{ij}^0$ for the network shown in (a). The rows and columns of the matrix were reordered after applying average linkage clustering, placing next to each other nodes with the highest topological overlap. The colors denote the degree of topological overlap between each node pair, as calculated in (a). By cutting the dendrogram with the orange line, it recovers the three modules built into the network. The dendrogram indicates that the EFG and the HIJK modules are closer to each other than they are to the ABC module.

The pairwise mergers of Step 3 will eventually pull all nodes into a single community. We can use a dendrogram to extract the underlying community organization.

The dendrogram visualizes the order in which the nodes are assigned to specific communities. For example, the dendrogram of Figure 9.9b tells us that the algorithm first merged nodes A with B, K with J and E with F, as each of these pairs have $x_{ij}=1$. Next node C was added to the (A, B) community, I to (K, J) and G to (E, F).

To identify the communities we must cut the dendrogram. Hierarchical clustering does not tell us where that cut should be. Using for example the cut indicated as a dashed line in Figure 9.9b, we recover the three obvious communities (ABC, EFG, and HIJK).

Applied to the E. coli metabolic network (Figure 9.3a), the Ravasz algorithm identifies the nested community structure of bacterial metabolism. To check the biological relevance of these communities, we color-coded the branches of the dendrogram according to the known biochemical classification of each metabolite. As shown in Figure 9.3b, substrates with similar biochemical role tend to be located on the same branch of the tree. In other words the known biochemical classification of these metabolites confirms the biological relevance of the communities extracted from the network topology.

**Computational Complexity**
How many computations do we need to run the Ravasz algorithm? The algorithm has four steps, each with its own computational complexity:

(a) **Similarity Matrix**
Seven nodes forming two distinct communities. The table shows the distance $r_{ij}$ between each node pair, acting as the similarity $x_{ij}$.

(b) **Single Linkage Clustering**
The similarity between communities 1 and 2 is the smallest of all $x_{ij}$, where $i$ and $j$ are in different communities. Hence the similarity is $x_{12}=1.59$, corresponding to the distance between nodes C and E.

(c) **Complete Linkage Clustering**
The similarity between two communities is the maximum of $x_{ij}$, where $i$ and $j$ are in different communities. Hence $x_{12}=3.97$.

(d) **Average Linkage Clustering**
The similarity between two communities is the average of $x_{ij}$ over all node pairs $i$ and $j$ that belong to different communities. This is the procedure implemented in the Ravasz algorithm, providing $x_{12}=2.84$. 

---

*Figure 9.10*
Cluster Similarity

In agglomerative clustering we need to determine the similarity of two communities from the node similarity matrix $x_{ij}$. We illustrate this procedure for a set of points whose similarity $x_{ij}$ is the physical distance $r_{ij}$ between them. In networks $x_{ij}$ corresponds to some network-based distance measure, like $x_{ij}$ defined in (9.7).

(a) **Similarity Matrix**
Seven nodes forming two distinct communities. The table shows the distance $r_{ij}$ between each node pair, acting as the similarity $x_{ij}$.

(b) **Single Linkage Clustering**
The similarity between communities 1 and 2 is the smallest of all $x_{ij}$, where $i$ and $j$ are in different communities. Hence the similarity is $x_{12}=1.59$, corresponding to the distance between nodes C and E.

(c) **Complete Linkage Clustering**
The similarity between two communities is the maximum of $x_{ij}$, where $i$ and $j$ are in different communities. Hence $x_{12}=3.97$.

(d) **Average Linkage Clustering**
The similarity between two communities is the average of $x_{ij}$ over all node pairs $i$ and $j$ that belong to different communities. This is the procedure implemented in the Ravasz algorithm, providing $x_{12}=2.84$. 

---

*Figure 9.3a*
E. coli metabolic network

*Figure 9.3b*
Biochemical classification of metabolites in the E. coli metabolic network.
Step 1: The calculation of the similarity matrix $x^0_{ij}$ requires us to compare $N^2$ node pairs, hence the number of computations scale as $N^2$. In other words its computational complexity is $O(N^2)$.

Step 2: Group similarity requires us to determine in each step the distance of the new cluster to all other clusters. Doing this $N$ times requires $O(N^2)$ calculations.

Steps 3 & 4: The construction of the dendrogram can be performed in $O(N \log N)$ steps.

Combining Steps 1-4, we find that the number of required computations scales as $O(N^2) + O(N^2) + O(N \log N)$. As the slowest step scales as $O(N^2)$, the algorithm’s computational complexity is $O(N^2)$. Hence hierarchal clustering is much faster than the brute force approach, which generally scales as $O(eN^2)$.

DIVISIVE PROCEDURES: THE GIRVAN-NEWMAN ALGORITHM

Divisive procedures systematically remove the links connecting nodes that belong to different communities, eventually breaking a network into isolated communities. We illustrate their use by introducing an algorithm proposed by Michelle Girvan and Mark Newman [9,23], consisting of the following steps:

Step 1: Define Centrality
While in agglomerative algorithms $x^0_y$ selects node pairs that belong to the same community, in divisive algorithms $x^0_y$, called centrality, selects node pairs that are in different communities. Hence we want $x^0_y$ to be high (or low) if nodes $i$ and $j$ belong to different communities and small if they are in the same community. Three centrality measures that satisfy this expectation are discussed in Figure 9.11. The fastest of the three is link betweenness, defining $x^0_y$ as the number of shortest paths that go through the link $(i,j)$. Links connecting different communities are expected to have large $x^0_y$ while links within a community have small $x^0_y$.

Step 2: Hierarchical Clustering
The final steps of a divisive algorithm mirror those we used in agglomerative clustering (Figure 9.12):

1. Compute the centrality $x^0_y$ of each link.
2. Remove the link with the largest centrality. In case of a tie, choose one link randomly.
3. Recalculate the centrality of each link for the altered network.

Figure 9.11
Centrality Measures

Divisive algorithms require a centrality measure that is high for nodes that belong to different communities and low for node pairs in the same community. Two frequently used measures can achieve this:

(a) Link Betweenness
Link betweenness captures the role of each link in information transfer. Hence $x^0_y$ is proportional to the number of shortest paths between all node pairs that run along the link $(i,j)$. Consequently, inter-community links, like the central link in the figure with $x^0_y=0.57$, have large betweenness. The calculation of link betweenness scales as $O(LN)$, or $O(N^2)$ for a sparse network [23].

(b) Random-Walk Betweenness
A pair of nodes $m$ and $n$ are chosen at random. A walker starts at $m$, following each adjacent link with equal probability until it reaches $n$. Random walk betweenness $x^0_y$ is the probability that the link $i \rightarrow j$ was crossed by the walker after averaging over all possible choices for the starting nodes $m$ and $n$. The calculation requires the inversion of an $N \times N$ matrix, with $O(N^3)$ computational complexity and averaging the flows over all node pairs, with $O(LN^2)$. Hence the total computational complexity of random walk betweenness is $O([L + N] N^2)$, or $O(N^3)$ for a sparse network.
Hierarchical clustering raises two fundamental questions:

(a) The divisive hierarchical algorithm of Girvan and Newman uses link betweenness (Figure 9.11a) as centrality. In the figure the link weights, assigned proportionally to \( x_{ij} \), indicate that links connecting different communities have the highest \( x_{ij} \). Indeed, each shortest path between these communities must run through them.

(b)-(d) The sequence of images illustrates how the algorithm removes one-by-one the three highest \( x_{ij} \) links, leaving three isolated communities behind. Note that betweenness needs to be recalculated after each link removal.

(e) The dendrogram generated by the Girvan-Newman algorithm. The cut at level 3, shown as an orange dotted line, reproduces the three communities present in the network.

(f) The modularity function, \( M \), introduced in Section 9.4, helps us select the optimal cut. Its maxima agrees with our expectation that the best cut is at level 3, as shown in (e).

4. Repeat steps 2 and 3 until all links are removed.

Girvan and Newman applied their algorithm to Zachary’s Karate Club (Figure 9.2a), finding that the predicted communities matched almost perfectly the two groups after the break-up. Only node 3 was classified incorrectly.

**Computational Complexity**

The rate limiting step of divisive algorithms is the calculation of centrality. Consequently the algorithm’s computational complexity depends on which centrality measure we use. The most efficient is link betweenness, with \( O(LN) \) [24,25,26] (Figure 9.11a). Step 3 of the algorithm introduces an additional factor \( L \) in the running time, hence the algorithm scales as \( O(L^2N) \), or \( O(N^2) \) for a sparse network.

**Hierarchical Clustering in Real Networks**

Hierarchical clustering raises two fundamental questions:

**Nested Communities**

First, it assumes that small modules are nested into larger ones. These nested communities are well captured by the dendrogram (Figures 9.9b and 9.12e). How do we know, however, if such hierarchy is indeed present in a network? Could this hierarchy be imposed by our algorithms, whether or not the underlying network has a nested community structure?

**Communities and the Scale-Free Property**

Second, the density hypothesis states that a network can be partitioned into a collection of subgraphs that are only weakly linked to other subgraphs. How can we have isolated communities in a scale-free network, if the hubs inevitably link multiple communities?
The hierarchical network model, whose construction is shown in Figure 9.13, resolves the conflict between communities and the scale-free property and offers intuition about the structure of nested hierarchical communities. The obtained network has several key characteristics:

**Scale-free Property**
The hierarchical model generates a scale-free network with degree exponent (Figure 9.14a, ADVANCED TOPICS 9.A)

\[ \gamma = 1 + \frac{\ln 5}{\ln 4} = 2.161. \]

**Size Independent Clustering Coefficient**
While for the Erdős-Rényi and the Barabási-Albert models the clustering coefficient decreases with N (SECTION 5.9), for the hierarchical network we have \( C = 0.743 \) independent of the network size (Figure 9.14c). Such N-independent clustering coefficient has been observed in metabolic networks [11].

**Hierarchical Modularity**
The model consists of numerous small communities that form larger communities, which again combine into ever larger communities. The quantitative signature of this nested hierarchical modularity is the dependence of a node's clustering coefficient on the node's degree [11,27,28]

\[ C(k) \sim k^{-1}. \]  

Equation (9.8) captures the way the communities are organized in a network. Indeed, small degree nodes have high C because they reside in dense communities. High degree nodes have small C because they connect to different communities. For example, in Figure 9.13c the nodes at the center of the five-node modules have \( k=4 \) and clustering coefficient \( C=4 \). Those at the center of a 25-node module have \( k=20 \) and \( C=3/19 \). Those at the center of the 125-node modules have \( k=84 \) and \( C=3/83 \). Hence the higher the degree of a node, the smaller is its C.

The hierarchical network model suggests that inspecting \( C(k) \) allows us to decide if a network is hierarchical. For the Erdős-Rényi and the Barabási-Albert models \( C(k) \) is independent of \( k \), indicating that they do not display hierarchical modularity. To see if hierarchical modularity is present in real systems, we calculated \( C(k) \) for ten reference networks, finding that (Figure 9.36):

- Only the power grid lacks hierarchical modularity, its \( C(k) \) being independent of \( k \) (Figure 9.36a).
- For the remaining nine networks \( C(k) \) decreases with \( k \). Hence in
these networks small nodes are part of small dense communities, while hubs link disparate communities to each other.

- For the scientific collaboration, metabolic, and citation network $C(k)$ follows (9.8) in the high-$k$ region. The form of $C(k)$ for the Internet, mobile, email, protein interactions, and the WWW needs to be derived individually, as for those $C(k)$ does not follow (9.8). More detailed network models predict $C(k) \sim k^\beta$, where $\beta$ is between 0 and 2 [27,28].

In summary, in principle hierarchical clustering does not require preliminary knowledge about the number and the size of communities. In practice it generates a dendrogram that offers a family of community partitions characterizing the studied network. This dendrogram does not tell us which partition captures best the underlying community structure. Indeed, any cut of the hierarchical tree offers a potentially valid partition (Figure 9.15). This is at odds with our expectation that in each network there is a ground truth, corresponding to a unique community structure.

Figure 9.15
Ambiguity in Hierarchical Clustering

Hierarchical clustering does not tell us where to cut a dendrogram. Indeed, depending on where we make the cut in the dendrogram of Figure 9.9a, we obtain (b) two, (c) three or (d) four communities. While for a small network we can visually decide which cut captures best the underlying community structure, it is impossible to do so in larger networks. In the next section we discuss modularity, that helps us select the optimal cut.
While there are multiple notions of hierarchy in networks [29,30], inspecting $C(k)$ helps decide if the underlying network has hierarchical modularity. We find that $C(k)$ decreases in most real networks, indicating that most real systems display hierarchical modularity. At the same time $C(k)$ is independent of $k$ for the Erdős-Rényi or Barabási-Albert models, indicating that these canonical models lack a hierarchical organization.
In a randomly wired network the connection pattern between the nodes is expected to be uniform, independent of the network’s degree distribution. Consequently these networks are not expected to display systematic local density fluctuations that we could interpret as communities. This expectation inspired the third hypothesis of community organization:

H3: Random Hypothesis

Randomly wired networks lack an inherent community structure.

This hypothesis has some actionable consequences: By comparing the link density of a community with the link density obtained for the same group of nodes for a randomly rewired network, we could decide if the original community corresponds to a dense subgraph, or its connectivity pattern emerged by chance.

In this section we show that systematic deviations from a random configuration allow us to define a quantity called modularity, that measures the quality of each partition. Hence modularity allows us to decide if a particular community partition is better than some other one. Finally, modularity optimization offers a novel approach to community detection.

**MODULARITY**

Consider a network with $N$ nodes and $L$ links and a partition into $n_c$ communities, each community having $N_c$ nodes connected to each other by $L_c$ links, where $c = 1,...,n_c$. If $L_c$ is larger than the expected number of links between the $N_c$ nodes given the network’s degree sequence, then the nodes of the subgraph $C_c$ could indeed be part of a true community, as expected based on the Density Hypothesis H2 (Figure 9.2). We therefore measure the difference between the network’s real wiring diagram ($A_{ij}$) and the expected number of links between $i$ and $j$ if the network is randomly wired ($p_{ij}$),

$$M_c = \frac{1}{2L} \sum_{(i,j) \in C_c} (A_{ij} - p_{ij}),$$

(9.9)
Here \( p_{ij} \) can be determined by randomizing the original network, while keeping the expected degree of each node unchanged. Using the degree preserving null model (7.1) we have

\[
p_{ij} = \frac{k_i k_j}{2L},
\]

(9.10)

If \( M_c \) is positive, then the subgraph \( C_c \) has more links than expected by chance, hence it represents a potential community. If \( M_c \) is zero then the connectivity between the \( N_c \) nodes is random, fully explained by the degree distribution. Finally, if \( M_c \) is negative, then the nodes of \( C_c \) do not form a community.

Using (9.10) we can derive a simpler form for the modularity (9.9) (ADVANCED TOPICS 9.B)

\[
M_c = \frac{L_c}{L} - \left( \frac{k_c}{2L} \right)^2,
\]

(9.11)

where \( L_c \) is the total number of links within the community \( C_c \) and \( k_c \) is the total degree of the nodes in this community.

To generalize these ideas to a full network consider the complete partition that breaks the network into \( n_c \) communities. To see if the local link density of the subgraphs defined by this partition differs from the expected density in a randomly wired network, we define the partition’s modularity by summing (9.11) over all \( n_c \) communities [23]

\[
M = \sum_{c=1}^{n_c} \left[ \frac{L_c}{L} - \left( \frac{k_c}{2L} \right)^2 \right].
\]

(9.12)

Modularity has several key properties:

• Higher Modularity Implies Better Partition
  The higher is \( M \) for a partition, the better is the corresponding community structure. Indeed, in Figure 9.16a the partition with the maximum modularity (\( M=0.41 \)) accurately captures the two obvious communities. A partition with a lower modularity clearly deviates from these communities (Figure 9.16b). Note that the modularity of a partition cannot exceed one [31,32].

• Zero and Negative Modularity
  By taking the whole network as a single community we obtain \( M=0 \), as in this case the two terms in the parenthesis of (9.12) are equal (Figure 9.16c). If each node belongs to a separate community, we have \( L_c=0 \) and the sum (9.12) has \( n_c \) negative terms, hence \( M \) is negative (Figure 9.16d).

We can use modularity to decide which of the many partitions predicted by a hierarchical method offers the best community structure, selecting the one for which \( M \) is maximal. This is illustrated in Figure 9.12f, which shows \( M \) for each cut of the dendrogram, finding a clear maximum when

Figure 9.16
Modularity
To better understand the meaning of modularity, we show \( M \) defined in (9.12) for several partitions of a network with two obvious communities.

(a) Optimal Partition
  The partition with maximal modularity \( M=0.41 \) closely matches the two distinct communities.

(b) Suboptimal Partition
  A partition with a sub-optimal but positive modularity, \( M=0.22 \), fails to correctly identify the communities present in the network.

(c) Single Community
  If we assign all nodes to the same community we obtain \( M=0 \), independent of the network structure.

(d) Negative Modularity
  If we assign each node to a different community, modularity is negative, obtaining \( M=-0.12 \).
the network breaks into three communities.

**THE GREEDY ALGORITHM**

The expectation that partitions with higher modularity corresponds to partitions that more accurately capture the underlying community structure prompts us to formulate our final hypothesis:

**H4: Maximal Modularity Hypothesis**

*For a given network the partition with maximum modularity corresponds to the optimal community structure.*

The hypothesis is supported by the inspection of small networks, for which the maximum $M$ agrees with the expected communities (Figures 9.12 and 9.16).

The maximum modularity hypothesis is the starting point of several community detection algorithms, each seeking the partition with the largest modularity. In principle we could identify the best partition by checking $M$ for all possible partitions, selecting the one for which $M$ is largest. Given, however, the exceptionally large number of partitions, this brute-force approach is computationally not feasible. Next we discuss an algorithm that finds partitions with close to maximal $M$, while bypassing the need to inspect all partitions.

**Greedy Algorithm**

The first modularity maximization algorithm, proposed by Newman [33], iteratively joins pairs of communities if the move increases the partition’s modularity. The algorithm follows these steps:

1. Assign each node to a community of its own, starting with $N$ communities of single nodes.
2. Inspect each community pair connected by at least one link and compute the modularity difference $\Delta M$ obtained if we merge them. Identify the community pair for which $\Delta M$ is the largest and merge them. Note that modularity is always calculated for the full network.
3. Repeat Step 2 until all nodes merge into a single community, recording $M$ for each step.
4. Select the partition for which $M$ is maximal.

To illustrate the predictive power of the greedy algorithm consider the collaboration network between physicists, consisting of $N=56,276$ scientists in all branches of physics who posted papers on arxiv.org (Figure 9.17). The greedy algorithm predicts about 600 communities with peak modularity $M = 0.713$. Four of these communities are very large, together containing 77% of all nodes (Figure 9.17a). In the largest community 93% of the authors publish in condensed matter physics while 87% of the authors in the second largest community publish in high energy physics.
physics, indicating that each community contains physicists of similar professional interests. The accuracy of the greedy algorithm is also illustrated in Figure 9.2a, showing that the community structure with the highest $M$ for the Zachary Karate Club accurately captures the club’s subsequent split.

**Computational Complexity**

Since the calculation of each $\Delta M$ can be done in constant time, Step 2 of the greedy algorithm requires $O(L)$ computations. After deciding which communities to merge, the update of the matrix can be done in a worst-case time $O(N)$. Since the algorithm requires $N-1$ community mergers, its complexity is $O((L + N)N)$, or $O(N^2)$ on a sparse graph. Optimized implementations reduce the algorithm’s complexity to $O(N \log^2 N)$ (ONLINE RESOURCE 9.1).

**LIMITS OF MODULARITY**

Given the important role modularity plays in community identification, we must be aware of some of its limitations.

**Resolution Limit**

Modularity maximization forces small communities into larger ones [34]. Indeed, if we merge communities A and B into a single community, the network’s modularity changes with (ADVANCED TOPICS 9.B)

$$\Delta M_{AB} = \frac{\ell_{AB}}{L} - \frac{k_A k_B}{2L^2}$$

where $\ell_{AB}$ is number of links that connect the nodes in community A with total degree $k_A$ to the nodes in community B with total degree $k_B$. If A and B are distinct communities, they should remain distinct when $M$ is maximized. As we show next, this is not always the case.

Consider the case when $k_A k_B/2L < 1$, in which case (9.13) predicts $\Delta M_{AB} > 0$ if there is at least one link between the two communities ($\ell_{AB} \geq 1$). Hence we must merge A and B to maximize modularity. Assuming for simplicity that $k_A = k_B = k$, if the total degree of the communities satisfies
then modularity increases by merging A and B into a single community, even if A and B are otherwise distinct communities. This is an artifact of modularity maximization: if \( k_A \) and \( k_B \) are under the threshold (9.14), the expected number of links between them is smaller than one. Hence even a single link between them will force the two communities together when we maximize \( M \). This resolution limit has several consequences:

- Modularity maximization cannot detect communities that are smaller than the resolution limit (9.14). For example, for the WWW sample with \( L = 1,497,134 \) (Table 2.1) modularity maximization will have difficulties resolving communities with total degree \( k_C < 1,730 \).

- Real networks contain numerous small communities [36-38]. Given the resolution limit (9.14), these small communities are systematically forced into larger communities, offering a misleading characterization of the underlying community structure.

To avoid the resolution limit we can further subdivide the large communities obtained by modularity optimization [33,34,39]. For example, treating the smaller of the two condensed-matter groups of Figure 9.17a as a separate network and feeding it again into the greedy algorithm, we obtain about 100 smaller communities with an increased modularity \( M = 0.807 \) (Figure 9.17b) [33].

**Modularity Maxima**

All algorithms based on maximal modularity rely on the assumption that a network with a clear community structure has an optimal partition with a maximal \( M \) [40]. In practice we hope that \( M_{\text{max}} \) is an easy to find maxima and that the communities predicted by all other partitions are distinguishable from those corresponding to \( M_{\text{max}} \). Yet, as we show next, this optimal partition is difficult to identify among a large number of close to optimal partitions.

Consider a network composed of \( n_C \) subgraphs with comparable link densities \( k_c = 2L/n_c \). The best partition should correspond to the one where each cluster is a separate community (Figure 9.18a), in which case \( M = 0.867 \). Yet, if we merge the neighboring cluster pairs into a single community we obtain a higher modularity \( M = 0.87 \) (Figure 9.18b). In general (9.13) and (9.14) predicts that if we merge a pair of clusters, we change modularity with

\[
\Delta M = \frac{L_{AB}}{L} - \frac{2}{n_c^2}.
\]

In other words the drop in modularity is less than \( \Delta M = -2/n_c^2 \). For a network with \( n_c = 20 \) communities, this change is at most \( \Delta M = -0.005 \), tiny compared to the maximal modularity \( M = 0.87 \) (Figure 9.18b). As the
In summary, modularity offers a first principle understanding of a network’s community structure. Indeed, (9.16) incorporates in a compact form a number of essential questions, like what we mean by a community, how we choose the appropriate null model, and how we measure the goodness of a particular partition. Consequently, modularity optimization plays a
central role in the community finding literature.

At the same time, modularity has several well-known limitations: First, it forces together small weakly connected communities. Second, networks lack a clear modularity maxima, developing instead a modularity plateau containing many partitions with hard to distinguish modularity. This plateau explains why numerous modularity maximization algorithms can rapidly identify a high $M$ partition: They identify one of the numerous partitions with close to optimal $M$. Finally, analytical calculations and numerical simulations indicate that even random networks contain high modularity partitions, at odds with the random hypothesis H3 that motivated the concept of modularity [41-43].

Modularity optimization is a special case of a larger problem: Finding communities by optimizing some quality function $Q$. The greedy algorithm and the Louvain algorithm described in ADVANCED TOPICS 9.C assume that $Q = M$, seeking partitions with maximal modularity. In ADVANCED TOPICS 9.C we also describe the Infomap algorithm, that finds communities by minimizing the map equation $L$, an entropy-based measure of the partition quality [44-46].
A node is rarely confined to a single community. Consider a scientist, who belongs to the community of scientists that share his professional interests. Yet, he also belongs to a community consisting of family members and relatives and perhaps another community of individuals sharing his hobby (Figure 9.19). Each of these communities consists of individuals who are members of several other communities, resulting in a complicated web of nested and overlapping communities [36]. Overlapping communities are not limited to social systems: The same genes are often implicated in multiple diseases, an indication that disease modules of different disorders overlap [14].

While the existence of a nested community structure has long been appreciated by sociologists [47] and by the engineering community interested in graph partitioning, the algorithms discussed so far force each node into a single community. A turning point was the work of Tamás Vicsek and collaborators [36,48], who proposed an algorithm to identify overlapping communities, bringing the problem to the attention of the network science community. In this section we discuss two algorithms to detect overlapping communities, clique percolation and link clustering.

**CLIQUE PERCOLATION**

The clique percolation algorithm, often called CFinder, views a community as the union of overlapping cliques [36]:

- Two $k$-cliques are considered adjacent if they share $k – 1$ nodes (Figure 9.20b).
- A $k$-clique community is the largest connected subgraph obtained by the union of all adjacent $k$-cliques (Figure 9.20c).
- $k$-cliques that can not be reached from a particular $k$-clique belong to other $k$-clique communities (Figure 9.20c,d).

The CFinder algorithm identifies all cliques and then builds an $N_{clique} \times N_{clique}$ clique–clique overlap matrix $O$, where $N_{clique}$ is the number of cliques and $O_{ij}$ is the number of nodes shared by cliques $i$ and $j$ (Figure 9.39). A typical...
output of the CFinder algorithm is shown in Figure 9.21, displaying the community structure of the word *bright*. In the network two words are linked to each other if they have a related meaning. We can easily check that the overlapping communities identified by the algorithm are meaningful: The word *bright* simultaneously belongs to a community containing light-related words, like *glow* or *dark*; to a community capturing colors (*yellow, brown*); to a community consisting of astronomical terms (*sun, ray*); and to a community linked to intelligence (*gifted, brilliant*). The example also illustrates the difficulty the earlier algorithms would have in identifying communities of this network: they would force *bright* into one of the four communities and remove from the other three. Hence communities would be stripped of a key member, leading to outcomes that are difficult to interpret.

Could the communities identified by CFinder emerge by chance? To distinguish the real *k*-clique communities from communities that are a pure consequence of high link density we explore the percolation properties of *k*-cliques in a random network [48]. As we discussed in Chapter 3, if a random network is sufficiently dense, it has numerous cliques of varying order. A large *k*-clique community emerges in a random network only if the connection probability *p* exceeds the threshold (Advanced Topics 9.D)

\[
p_c(k) = \frac{1}{[(k-1)N]^{1/(k-1)}}.
\]  

(9.16)

Under *p_c*(k) we expect only a few isolated *k*-cliques (Figure 9.22a). Once *p* exceeds *p_c*(k), we observe numerous cliques that form *k*-clique communities (Figure 9.22b). In other words, each *k*-clique community has its own threshold:

- For *k*=2 the *k*-cliques are links and (9.16) reduces to *p_c*(k)=1/N, which
is the condition for the emergence of a giant connected component in Erdős–Rényi networks.

- For \( k=3 \) the cliques are triangles (Figure 9.22a,b) and (9.16) predicts \( p_c(k) \approx 1/\sqrt{2N} \).

In other words, \( k \)-clique communities naturally emerge in sufficiently dense networks. Consequently, to interpret the overlapping community structure of a network, we must compare it to the community structure obtained for the degree-randomized version of the original network.

**Computational Complexity**

Finding cliques in a network requires algorithms whose running time grows exponentially with \( N \). Yet, the CFinder community definition is based on cliques instead of maximal cliques, which can be identified in polynomial time [49]. If, however, there are large cliques in the network, it is more efficient to identify all cliques using an algorithm with \( O(eN) \) complexity [36]. Despite this high computational complexity, the algorithm is relatively fast, processing the mobile call network of 4 million mobile phone users in less than one day [50] (see also Figure 9.28).

**LINK CLUSTERING**

While nodes often belong to multiple communities, links tend to be community specific, capturing the precise relationship that defines a node’s membership in a community. For example, a link between two individuals may indicate that they are in the same family, or that they work together, or that they share a hobby, designations that only rarely overlap. Similarly, in biology each binding interaction of a protein is responsible for a different function, uniquely defining the role of the protein in the cell. This specificity of links has inspired the development of community finding algorithms that cluster links rather than nodes [51,52].

The link clustering algorithm proposed by Ahn, Bagrow and Lehmann [51] consists of the following steps:

**Step 1: Define Link Similarity**

The similarity of a link pair is determined by the neighborhood of the nodes connected by them. Consider for example the links \((i,k)\) and \((j,k)\), connected to the same node \(k\). Their similarity is defined as (Figure 9.23a-c)

\[
S((i,k),(j,k)) = \frac{|n_i(i) \cap n_j(j)|}{|n_i(i) \cup n_j(j)|}, \quad (9.17)
\]

where \( n_i(i) \) is the list of the neighbors of node \( i \), including itself. Hence \( S \) measures the relative number of common neighbors \( i \) and \( j \) have. Consequently \( S=1 \) if \( i \) and \( j \) have the same neighbors (Figure 9.23c). The less is the overlap between the neighborhood of the two links, the smaller is \( S \) (Figure 9.23b).
Step 2: Apply Hierarchical Clustering

The similarity matrix $S$ allows us to use hierarchical clustering to identify link communities (SECTION 9.3). We use a single-linkage procedure, iteratively merging communities with the largest similarity link pairs (Figure 9.10).

Taken together, for the network of Figure 9.23e, (9.17) provides the similarity matrix shown in (d). The single-linkage hierarchical clustering leads to the dendrogram shown in (d), whose cuts result in the link communities shown in (e) and the overlapping node communities shown in (f).

Figure 9.24 illustrates the community structure of the characters of Victor Hugo’s novel *Les Miserables* identified using the link clustering algorithm. Anyone familiar with the novel can convince themselves that the communities accurately represent the role of each character. Several characters are placed in multiple communities, reflecting their overlapping roles in the novel. Links, however, are unique to each community.

Computational Complexity

The link clustering algorithm involves two time-limiting steps: similarity calculation and hierarchical clustering. Calculating the similarity (9.17) for a link pair with degrees $k_i$ and $k_j$ requires $\max(k_i, k_j)$ steps. For a scale-free network with degree exponent $\gamma$ the calculation of similarity has complexity $O(N^{2/\gamma - 1})$, determined by the size of the largest node, $k_{max}$. Hierarchical clustering requires $O(L^2)$ time steps. Hence the algorithm’s total complexity is $O(N^{2/\gamma - 1} + L^2)$. 

(a) The similarity $S$ of the $(i,k)$ and $(j,k)$ links connected to node $k$ detects if the two links belong to the same group of nodes. Denoting with $n_i(i)$ the list of neighbors of node $i$, including itself, we obtain $|n_i(i) \cap n_j(j)| = 12$ and $|n_i(i) \cap n_j(j)| = 4$, resulting in $S = 1/3$ according to (9.17).

(b) For an isolated ($k_i = k_j = 1$) connected triple we obtain $S = 1/3$.

(c) For a triangle we have $S = 1$.

(d) The link similarity matrix for the network shown in (e) and (f). Darker entries correspond to link pairs with higher similarity $S$. The figure also shows the resulting link dendrogram.

(e) The link community structure predicted by the cut of the dendrogram shown as an orange dashed line in (d).

(f) The overlapping node communities derived from the link communities shown in (e).

After [51].
The computational complexity is $O(N^2/(\gamma-1)) + O(L^2)$. For sparse graphs the latter term dominates, leading to $O(N^2)$.

The need to detect overlapping communities have inspired numerous algorithms [53]. For example, the CFinder algorithm has been extended to the analysis of weighted [54], directed and bipartite graphs [55,56]. Similarly, one can derive quality functions for link clustering [52], like the modularity function discussed in Section 9.4.

In summary, the algorithms discussed in this section acknowledge the fact that nodes naturally belong to multiple communities. Therefore by forcing each node into a single community, as we did in the previous sections, we obtain a misleading characterization of the underlying community structure. Link communities recognize the fact that each link accurately captures the nature of the relationship between two nodes. As a bonus link clustering also predicts the overlapping community structure of a network.
Community identification algorithms offer a powerful diagnosis tool, allowing us to characterize the local structure of real networks. Yet, to interpret and use the predicted communities, we must understand the accuracy of our algorithms. Similarly, the need to diagnose large networks prompts us to address the computational efficiency of our algorithms. In this section we focus on the concepts needed to assess the accuracy and the speed of community finding.

**ACCURACY**

If the community structure is uniquely encoded in the network’s wiring diagram, each algorithm should predict precisely the same communities. Yet, given the different hypotheses the various algorithms embody, the partitions uncovered by them can differ, prompting the question: Which community finding algorithm should we use?

To assess the performance of community finding algorithms we need to measure an algorithm’s accuracy, i.e. its ability to uncover communities in networks whose community structure is known. We start by discussing two benchmarks, which are networks with predefined community structure, that we can use to test the accuracy of a community finding algorithm.

**Girvan-Newman (GN) Benchmark**

The Girvan-Newman benchmark consists of $N=128$ nodes partitioned into $n_c=4$ communities of size $N_c=32$ [9,57]. Each node is connected with probability $p_{int}$ to the $N_c-1$ nodes in its community and with probability $p_{ext}$ to the $3N_c$ nodes in the other three communities. The control parameter

$$
\mu = \frac{k_{ext}}{k_{ext} + k_{int}},
$$

(9.18)

captures the density differences within and between communities. We
In Figure 9.25, the position of each node in (a) and (c) shows the planted communities of the Girvan-Newman (GN) benchmark, illustrating the presence of four distinct communities, each with \( N_c = 32 \) nodes.

(a) The node colors represent the partitions predicted by the Ravasz algorithm for mixing parameter \( \mu = 0.40 \) given by (9.18). As in this case the communities are well separated, we have an excellent agreement between the planted and the detected communities.

(b) The normalized mutual information in function of the mixing parameter \( \mu \) for the Ravasz algorithm. For small \( \mu \) we have \( I_n \approx 1 \) and \( n_c \approx 4 \), indicating that the algorithm can easily detect well separated communities, as illustrated in (a). As we increase \( \mu \) the link density difference within and between communities becomes less pronounced. Consequently the communities are increasingly difficult to identify and \( I_n \) decreases.

(c) For \( \mu = 0.50 \) the Ravasz algorithm misplaces a notable fraction of the nodes, as in this case the communities are not well separated, making it harder to identify the correct community structure.

Note that the Ravasz algorithm generates multiple partitions, hence for each \( \mu \) we show the partition with the largest modularity, \( M \). Next to (a) and (c) we show the normalized mutual information associated with the corresponding partition and the number of detected communities \( n_c \). The normalized mutual information (9.23), developed for non-overlapping communities, can be extended to overlapping communities as well [59].

Expect community finding algorithms to perform well for small \( \mu \) (Figure 9.25a), when the probability of connecting to nodes within the same community exceeds the probability of connecting to nodes in different communities. The performance of all algorithms should drop for large \( \mu \) (Figure 9.25b), when the link density within the communities becomes comparable to the link density in the rest of the network.

Lancichinetti-Fortunato-Radicchi (LFR) Benchmark
The GN benchmark generates a random graph in which all nodes have comparable degree and all communities have identical size. Yet, the degree distribution of most real networks is fat tailed, and so is the community size distribution (Figure 9.29). Hence an algorithm that performs well on the GN benchmark may not do well on real networks. To avoid
this limitation, the LFR benchmark (Figure 9.26) builds networks for which both the node degrees and the planted community sizes follow power laws [58].

Having built networks with known community structure, next we need tools to measure the accuracy of the partition predicted by a particular community finding algorithm. As we do so, we must keep in mind that the two benchmarks discussed above correspond to a particular definition of communities. Consequently algorithms based on clique percolation or link clustering, that embody a different notion of communities, may not fare so well on these.

**Measuring Accuracy**

To compare the predicted communities with those planted in the benchmark, consider an arbitrary partition into non-overlapping communities. In each step we randomly choose a node and record the label of the community it belongs to. The result is a random string of community labels that follow a $p(C)$ distribution, representing the probability that a randomly selected node belongs to the community $C$.

Consider two partitions of the same network, one being the benchmark (ground truth) and the other the partition predicted by a community finding algorithm. Each partition has its own $p(C_1)$ and $p(C_2)$ distribution. The joint distribution, $p(C_1, C_2)$, is the probability that a randomly chosen node belongs to community $C_1$ in the first partition and $C_2$ in the second. The similarity of the two partitions is captured by the normalized mutual information [38]

$$I_n = \frac{\sum_{C_1, C_2} p(C_1, C_2) \log_2 \frac{p(C_1, C_2)}{p(C_1)p(C_2)}}{\frac{1}{2}H(p(C_1)) + \frac{1}{2}H(p(C_2))}. \quad (9.19)$$

The numerator of (9.19) is the **mutual information** $I$, measuring the information shared by the two community assignments: $I=0$ if $C_1$ and $C_2$ are independent of each other; $I$ equals the maximal value $H(p(C_1)) = H(p(C_2))$ when the two partitions are identical and

$$H(p(C)) = - \sum_C p(C) \log_2 p(C) \quad (9.20)$$
is the Shannon entropy.

If all nodes belong to the same community, then we are certain about the next label and $H=0$, as we do not gain new information by inspecting the community to which the next node belongs to. $H$ is maximal if $p(C)$ is the uniform distribution, as in this case we have no idea which community comes next and each new node provides $H$ bits of new information.

In summary, $I_n=1$ if the benchmark and the detected partitions are identical, and $I_n=0$ if they are independent of each other. The utility of $I_n$ is illustrated in Figure 9.25b that shows the accuracy of the Ravasz algorithm for the Girvan-Newman benchmark. In Figure 9.27 we use $I_n$ to test the performance of each algorithm against the GN and LFR benchmarks. The results allow us to draw several conclusions:

- We have $I_n=1$ for $\mu<0.5$. Consequently when the link density within communities is high compared to their surroundings, most algorithms accurately identify the planted communities. Beyond $\mu=0.5$ the accuracy of each algorithm drops.

- The accuracy is benchmarks dependent. For the more realistic LFR benchmark the Louvain and the Ravasz methods offer the best performance and greedy modularity performs poorly.

SPEED

As discussed in Section 9.2, the number of possible partitions increases faster than exponentially with $N$, becoming astronomically high for most real networks. While community identification algorithms do not check all partitions, their computational cost still varies widely, determining their speed and consequently the size of the network they can handle. Table 9.1 summarizes the computational complexity of the algorithms discussed in this chapter. Accordingly, the most efficient are the Louvain and the Infomap algorithms, both of which scale as $O(N\log N)$. The least efficient is CFinder with $O(e^N)$.

These scaling laws do not capture the actual running time, however. They only show how the running time scales with $N$. This scaling matters if we need to find communities in very large networks. To get a sense of the true speed of these algorithms we measured their running time for the protein interaction network ($N=2,018$), the power grid ($N=4,941$) and the

---

**Figure 9.27**

Testing Against Benchmarks

We tested each community finding algorithm that predicts non-overlapping communities against the GN and the LFR benchmarks. The plots show the normalized mutual information $I_n$ against $\mu$ for five algorithms. For the naming of each algorithm, see Table 9.1.

(a) GN Benchmark
The horizontal axis shows the mixing parameter (9.18), representing the fraction of links connecting different communities. The vertical axis is the normalized mutual information (9.19). Each curve is averaged over 100 independent realizations.

(b) LFR Benchmark
Same as in (a) but for the LFR benchmark. The benchmark parameters are $N=1,000$, $\langle k \rangle=20$, $\gamma=2$, $k_{\text{max}}=50$, $\zeta=1$, maximum community size: 100, minimum community size: 20. Each curve is averaged over 25 independent realizations.
The computational complexity of the community identification algorithms discussed in this chapter. While computational complexity depends on both N and L, for sparse networks with good approximation we have L~N. We therefore list computational complexity in terms of N only.

![Graph](image_url)
collaboration network in seven days.

In summary, benchmarks allow us to compare the accuracy and the speed of the available algorithms. Given that the development of the fastest and the most accurate community detection tool remains an active arms race, those interested in the subject should consult the literature that compares algorithms across multiple dimensions [31, 58, 60, 61].
Research in network science is driven by the desire to quantify the fundamental principles that govern how networks emerge and how they are organized. These organizing principles impact the structure of communities, as well as our ability to identify them. In this section we discuss community evolution, the characteristics of community size distribution and the role of the link weights in community identification, allowing us to uncover the generic principles of community organization.

COMMUNITY SIZE DISTRIBUTION

According to the fundamental hypothesis (H1) the number and the size of communities in a network are uniquely determined by the network’s wiring diagram. We must therefore ask: What is the size distribution of these communities?

Many studies report fat tailed community size distributions, implying that numerous small communities coexist with a few very large ones [16,33,35,36,60]. To explore how widespread this pattern is, in Figure 9.29 we show $p_{N_c}$ for three networks, as predicted by various community finding algorithms. The plots indicate several patterns:

- For the protein interaction and the science collaboration network all algorithms predict an approximately fat tailed $p_{N_c}$. Hence in these networks numerous tiny communities coexist with a few large communities.

- For the power grid different algorithms lead to distinct outcomes. Modularity-based algorithms predict communities with comparable size $N_c = 10^2$. In contrast, the Ravasz algorithm and Infomap predict numerous communities with size $N_c = 10$ and a few larger communities. Finally, clique percolation and link clustering predict an approximately fat tailed community size distribution.

These differences suggest that the fat tailed community size distribution is not a byproduct of a particular algorithm. Rather it is an inherent...
property of some networks, like the protein and the scientific collaboration network. The different outcomes for the power grid suggests that this network lacks a unique and detectable community structure.

**COMMUNITIES AND LINK WEIGHTS**

Link weights are deeply correlated with the community structure. Yet, as we discuss next, the nature of these correlations is system dependent.

**Social Networks**

The more time two individuals spend together, the more likely that they share friends, which increases the chance that they belong to the same community. Consequently communities in social networks tend to be nucleated around strong ties. Links connecting different communities are weaker in comparison. This pattern, known as the weak tie hypothesis [62], is illustrated in Figure 9.30a for the mobile call network [63]. We observe that strong ties are indeed predominantly within the numerous small communities, and links connecting communities are visibly weaker.

**Transport Systems**

The purpose of many technological and biological networks is to transport materials or information. In this case the link weights are expected to correlate with betweenness centrality [64,65,66], a proxy of the local traffic carried by a network. As links connecting different communities must transport considerable amount of traffic, in transport networks strong ties are between communities. In contrast links within communities are weaker in comparison (Figure 9.30b).

The coupling between link weights and community structure suggests that incorporating the link weights could enhance the accuracy of community finding algorithms. Yet, the different nature of the coupling in social and technological systems serves as a cautionary note: Algorithms that aim to place in the same community nodes connected by strong ties may be only effective in social systems. They may offer potentially misleading results in technological and biological systems, where strong ties connect different communities.

**COMMUNITY EVOLUTION**

Changes in a network’s wiring diagram can have multiple consequences for communities: they can lead to the birth of new communities, the growth or the contraction of the existing communities, communities can merge with each other or split into several smaller communities, and finally communities can die (Figure 9.31) [50]. Studies focusing on social and communication networks offer several insights into the changes communities experience [50,67-73]:

**Growth**

The mobile call network helps us illustrate the relationship between link weights and communities. Links represent mutual calls between the users. We show only nodes that are at distance six or less from the individual highlighted as a black circle in (a).

(a) Real Weights

The link colors capture the aggregate call duration in minutes (see color bar). In line with the weak tie hypothesis we find strong ties mainly within communities and weak ties between communities [62].

(b) Betweenness Centrality

If the link weights are driven by the need to transport information or materials, as it is often the case in technological and biological systems, the weights are well approximated by betweenness centrality (Figure 9.11). We colored the links based on each link’s betweenness centrality. As the figure indicates, links connecting communities have high betweenness (red), whereas the links within communities have low betweenness (green).

After [63].
The probability that a node joins a community grows with the number of links the node has to members of that community [73].

**Contraction**
Nodes with only a few links to members of their community are more likely to leave the community than nodes with multiple links to community members [73]. In weighted networks the probability that a node leaves a community increases with the sum of its link weights to nodes outside the community.

**Splitting or Death**
The probability that a community disintegrates increases with the aggregate link weights to nodes outside the community.

**Age**
There is positive correlation between the age of a community and its size, indicating that older communities tend to be larger [50].

**Community Stability**
The membership of large communities changes faster with time than the membership of smaller communities. Indeed, in social networks large communities often correspond to institutions, companies or schools, that renew themselves by accepting new members, hiring new employees or enrolling new students. For small communities stability requires stable membership [50].

These results were obtained in the context of social systems. Our understanding of the patterns that cover community evolution in technological or biological systems remains limited.

In summary, several recurring patterns characterize the organization and the evolution of communities. The community size distribution is typically fat tailed, indicating the coexistence of many small communities with a few large ones. We also find system-dependent correlations between the community structure and link weights, so that in social systems the strong ties are mainly within communities, while in transport systems they are between communities. Finally, we gained an increasing understanding of the dynamical patterns that govern community evolution.

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**Figure 9.31**

**Evolving Communities**

When networks evolve in time, so does the underlying community structure. All changes in community structure are the result of six elementary events in the life of a community, illustrated in the figure: a community can grow or contract; communities can merge or may split; new communities are born while others may disappear. After [50].
The ubiquity of communities across different networks has turned community identification into a dynamically developing chapter of network science. Many of the developed algorithms are now available as software packages, allowing their immediate use for network diagnosis. Yet, the efficient use of these algorithms and the interpretation of their predictions requires us to be aware of the assumptions built into them. In this chapter we provided the intellectual and the quantitative foundations of community detection, helping us understand the origin and the assumptions behind the most frequently used algorithms.

Despite the successes of community identification, the field is faced with numerous open questions:

**Do We Really Have Communities?**
Throughout this chapter we avoided a fundamental question: How do we know that there are indeed communities in a particular network? In other words, could we decide that a network has communities without first identifying the communities themselves? The lack of an answer to this question represents perhaps the most glaring gap of the community finding literature. Community finding algorithms are designed to identify communities, whether they are there or not.

**Hypotheses or Theorems?**
Community identification relies on four hypotheses, summarized in BOX 9.3. We call them hypotheses because we can not prove their correctness. Further advances might be able to turn the Fundamental, the Random and the Maximal Modularity Hypotheses into theorems. Or we may learn about their limitations, as we did in the case of the Maximal Modularity Hypothesis (SECTION 9.6).

**Must all Nodes Belong to Communities?**
Community detection algorithm force all nodes into communities. This is likely an overkill for most real networks: some nodes belong to a single community, others to multiple communities, and likely many nodes

### BOX 9.3

**AT A GLANCE: COMMUNITIES**
Community identification rests on several hypotheses, pertaining to the nature of communities:

**Fundamental Hypothesis**
Communities are uniquely encoded in a network’s wiring diagram. They represent a grand truth that remains to be discovered using appropriate algorithms.

**Connectedness and Density Hypothesis**
A community corresponds to a locally dense connected subgraph.

**Random Hypothesis**
Randomly wired networks do not have communities.

**Maximal Modularity Hypothesis**
The partition with the maximum modularity offers the best community structure, where modularity is given by

\[
M = \sum_{c=1}^{L} \left( I - \left( \frac{k_c}{2L} \right)^2 \right)
\]
do not belong to any community. Most algorithms used in community identification do not make this distinction, forcing instead all nodes into some community.

**Dense vs. Sparse Communities**

Most networks explored in this book are sparse. Yet, with improvements in data collection, many real network maps will likely gain numerous links. In dense networks we often see numerous highly overlapping communities, forcing us to reevaluate the validity of the various hypotheses, and the appropriateness of the community detection algorithms discussed in this chapter. For example, in highly overlapping communities nodes may have higher external than internal degrees, limiting the validity of the density hypothesis.

**Do Communities Matter?**

We resort to an example to answer this question. Figure 9.32a shows a local neighborhood of the mobile call network, highlighting four communities identified by the link clustering algorithm (SECTION 9.5). The figure also shows the call frequency at noon (b) and at midnight (c), documenting different calling habits at different parts of the day. We find that the members of the top right community, shown as brown nodes in (a), are active at midnight (b), but they stop calling each other at noon (c). In contrast the light and the dark blue communities are active at noon, but are sleepy at midnight. This indicates that communities, identified only from the network’s wiring diagram, have coherent community-specific activity patterns.

Figure 9.32 suggests that once present, communities have a profound impact on network behavior. Numerous measurements support this conclusion: Information travels fast within a community but has difficulty reaching other communities [63]; communities influence the link weights [62]; the presence of communities can lead to degree correlations [74].

Communities are equally remarkable for their potential applications. For example, strengthening the links between clients that belong to the same community on the WWW can improve the performance of Web-based services [75]. In marketing, community finding can be used to identify customers with similar interests or purchasing habits, helping design efficient product recommendation systems [76]. Communities are often used to create data structures that can handle queries in a timely fashion [77,78]. Finally, community finding algorithms run behind many social networks sites, like Facebook, Twitter, or LinkedIn, helping these services discover potential friends, posts of interests and target advertising.

While community finding has deep roots in social and computer science, it is a relatively young chapter of network science (BOX 9.4). As such, our understanding of community organization continues to develop rapidly, offering increasingly accurate tools to diagnose the local structure of large networks.
Box 9.4

Community Finding: A Brief History

Early Communities
George Homans recorded the communication of bank tellers (top), identifying their communities (bottom) [3].

Graph Partitioning
Predecessors to community finding, graph partitioning algorithms optimize the layout of integrated circuits.

Michelle Girvan and Mark Newman propose the hierarchical divisive algorithm, igniting an explosive interest in community identification [9]. They also introduce modularity in 2004 [23].

Erzsébet Ravasz proposes a hierarchical agglomerative algorithm, unleashing an explosion of research within systems biology [11].

The Roots
Many of the concepts used in community finding have their roots in social and computer science, preceding network science.

The Explosion
The current interest in community finding was ignited by two papers, proposing algorithms to identify communities in social [9] and [11] biological systems.
9.1. Hierarchical Networks

Calculate the degree exponent of the hierarchical network shown in Figure 9.33.

9.2. Communities on a Circle

Consider a one dimensional lattice with \( N \) nodes that form a circle, where each node connects to its two neighbors. Partition the line into \( n_c \) consecutive clusters of size \( N_c = N/n_c \).

(a) Calculate the modularity of the obtained partition.
(b) According to the Maximum Modularity Hypothesis (SECTION 9.4), the maximum of \( M_c \) corresponds to the best partition. Obtain the community size \( n_c \) corresponding to the best partition.

9.3. Modularity Resolution Limit

Consider a network consisting of a ring of \( n_c \) cliques, each clique having \( N_c \) nodes and \( m(m-1)/2 \) links. The neighboring cliques are connected by a single link (Figure 9.34). The network has an obvious community structure, each community corresponding to a clique.

(a) Determine the modularity \( M_{\text{single}} \) of this natural partition, and the modularity \( M_{\text{pairs}} \) of the partition in which pairs of neighboring cliques are merged into a single community, as indicated by the dotted lines in Figure 9.34.
(b) Show that only for \( n_c < 2L \) will the modularity maximum predict the intuitively correct community partition, where

\[
L = n_c m(m-1) / 2 + n_c .
\]

(c) Discuss the consequences of violating the above inequality.


Show that the maximum value of modularity \( M \) defined in (9.12) cannot exceed one.
In this section we discuss the scaling properties of the hierarchical model introduced in Figure 9.13. We calculate the degree distribution and the degree-dependent clustering coefficient, deriving (9.8). Finally, we explore the presence of hierarchy in the ten real networks.

**Degree Distribution**

To compute the model’s degree distribution we count the nodes with different degrees. Starting with the five nodes of the first module in Figure 9.13a, we label the middle one a hub and call the remaining four nodes peripheral. All copies of this hub are again called hubs and we continue calling copies of peripheral nodes peripheral (Figure 9.35).

The largest hub at the center of the network acquires \( 4^n \) links during the \( n \)th iteration. Let us call this central hub \( H_n \) and the four copies of this hub \( H_{n-1} \) (Figure 9.35). We call \( H_{n-2} \) the \( 4 \times 5 \) leftover module centers whose size equals the size of the network at the \( (n-2) \)th iteration.

At the \( n \)th iteration the degree of the hub \( H_i \) follows

\[
k_n(H_i) = \sum_{l=1}^{i} 4^l = \frac{4}{3} \left( 4^i - 1 \right),
\]

where we used

\[
\sum_{l=0}^{i} x^l = \frac{x^{i+1} - 1}{x - 1},
\]

or

\[
\sum_{l=1}^{i} x^l = \frac{x^{i+1} - 1}{x - 1} - 1.
\]

For \( i < n \) the number of \( H_i \) modules is

\[
N_n(H_i) = 4 \times 5^{n-i-1}.
\]
i.e. there are four modules for $i = n - 1$; $4 \cdot 5$ modules for $i = n - 2$; ...; and $4 \cdot 5^{n-2}$ for $i=n$. Since we have $4 \cdot 5^{n-1}$ $H_i$-type hubs of degree $k_n(H_i)$, (4.35) and (4.38) allow us to write

$$\ln N_n(H_i) = C_n - i \cdot \ln 5 \quad (9.39)$$

$$\ln k_n(H_i) \approx i \cdot \ln 4 + \ln (4/3) \quad (9.40)$$

where

$$C_n = \ln 4 + (n - 1) \ln 5. \quad (9.41)$$

Note that in (9.40) we used the approximation $4^{i-1} \approx 4^i$.

For all $k > n + 2$ we can combine (9.40) and (9.41) to obtain

$$\ln N_n(H_i) = C_n' - \ln k_i \frac{\ln 5}{\ln 4} \quad (9.42)$$

or

$$N_n(H_i) \sim k_i^{-\frac{\ln 5}{\ln 4}}. \quad (9.43)$$

To calculate the degree distribution we need to normalize $N_n(H_i)$ by calculating the ratio

$$p_i \sim \frac{N_n(H_i)}{k_{i+1} - k_i} \sim k_i^{-\gamma}. \quad (9.44)$$

Using

$$k_{i+1} - k_i = \sum_{l=1}^{i+1} 4^l - \sum_{l=1}^{i} 4^l = 4^{i+1} = 3k_i + 4 \quad (9.45)$$

we obtain

$$p_i = \frac{k_i^{-\frac{\ln 5}{\ln 4}}}{3k_i + 4} \sim k_i^{-1 - \frac{\ln 4}{\ln 5}}. \quad (9.46)$$

In other words the obtained hierarchical network’s degree exponent is

$$\gamma = 1 + \frac{\ln 5}{\ln 4} = 2.16. \quad (9.47)$$

**Clustering Coefficient**

It is somewhat straightforward to calculate the clustering coefficient of the $H_i$ hubs. Their $\sum_{l=1}^{i} 4^l$ links come from nodes linked in a square, thus the connections between them equals their number. Consequently the number of links between the $H_i$’s neighbors is

$$\sum_{l=1}^{i} 4^l = k_n(H_i), \quad (9.48)$$

providing

$$C(H_i) = \frac{2k_i}{k_i(k_i - 1)} = \frac{2}{k_i - 1}. \quad (9.49)$$

In other words we obtain
indicating that $C(k)$ for the hubs scales as $k^{-1}$, in line with \ref{eq:chi}.

\begin{equation}
C(k) \approx \frac{2}{k}, \tag{9.50}
\end{equation}

\textbf{Empirical Results}

Figure 9.36 shows the $C(k)$ function for the ten reference networks. We also show $C(k)$ for each network after we applied degree-preserving randomization (green symbols), allowing us to make several observations:

- For small $k$ all networks have an order of magnitude higher $C(k)$ than their randomized counterpart. Therefore the small degree nodes are located in much denser neighborhoods than expected by chance.

- For the scientific collaboration, metabolic, and citation networks with a good approximation we have $C(k) \sim k^{-1}$, while the randomized $C(k)$ is flat. Hence these networks display the hierarchical modularity of the model of Figure 9.13.

- For the Internet, mobile phone calls, actors, email, protein interactions and the WWW $C(k)$ decreases with $k$, while their randomized $C(k)$ is $k$-independent. Hence while these networks display a hierarchical modularity, the observed $C(k)$ is not captured by our simple hierarchical model. To fit the $C(k)$ of these systems we need to build models that accurately capture their evolution. Such models predict that $C(k) \sim k^{-\beta}$, where $\beta$ can be different from one \cite{27}.

- Only for the power grid we observe a flat, $k$-independent $C(k)$, indicating the lack of a hierarchical modularity.

Taken together, Figure 9.36 indicates that most real networks display some nontrivial hierarchical modularity.
Figure 9.36
Hierarchy in Real Networks

The scaling of $C(k)$ with $k$ for the ten reference networks (purple symbols). The green symbols show $C(k)$ obtained after applying degree preserving randomization to each network, that washes out the local density fluctuations. Consequently communities and the underlying hierarchy are gone. Directed networks were made undirected to measure $C(k)$. The dashed line in each figure has slope -1, following (9.8), serving as a guide to the eye.
In this section we derive the expressions (9.12) and (9.13), characterizing the modularity function and its changes.

**Modularity as a Sum Over Communities**

Using (9.9) and (9.10) we can write the modularity of a full network as

\[
M = \frac{1}{2L} \sum_{i,j=1}^{N} (A_{ij} - \frac{k_i k_j}{2L}) \delta_{C_i, C_j},
\]

where \(C_i\) is the label of the community to which node \(i\) belongs to. As only node pairs that belong to the same community contribute to the sum in (9.51), we can rewrite the first term as a sum over communities,

\[
\frac{1}{2L} \sum_{i,j=1}^{N} A_{ij} \delta_{C_i, C_j} = \frac{1}{2L} \sum_{c=1}^{n_c} \sum_{i,j \in C_c} A_{ij} = \sum_{c=1}^{n_c} \frac{L_c}{L}
\]

where \(L_c\) is the number of links within community \(C_c\). The factor 2 disappears because each link is counted twice in \(A_{ij}\).

In a similar fashion the second term of (9.51) becomes

\[
\frac{1}{2L} \sum_{i,j=1}^{N} k_i k_j \delta_{C_i, C_j} = \frac{1}{2L} \sum_{c=1}^{n_c} \frac{1}{(2L)^2} \sum_{i,j \in C_c} k_i k_j = \sum_{c=1}^{n_c} \frac{k_c^2}{4L^2},
\]

where \(k_c\) is the total degree of the nodes in community \(C_c\). Indeed, in the configuration model the probability that a stub connects to a randomly chosen stub is \(\frac{1}{2L}\), as in total we have \(2L\) stubs in the network. Hence the likelihood that our stub connects to a stub inside the module is \(\frac{k_c}{2L}\). By repeating this procedure for all \(k_c\) stubs within the community \(C_c\) and adding 1/2 to avoid double counting, we obtain the last term of (9.53).

Combining (9.52) and (9.53) leads to (9.12).
Merging Two Communities

Consider communities A and B and denote with $k_A$ and $k_B$ the total degree in these communities (equivalent with $k_c$ above). We wish to calculate the change in modularity after we merge these two communities. Using (9.12), this change can be written as

$$
\Delta M_{AB} = \left[ \frac{L_{AB}}{L} - \left( \frac{k_{AB}}{2L} \right)^2 \right] - \left[ \frac{L_A}{L} - \left( \frac{k_A}{2L} \right)^2 + \frac{L_B}{L} - \left( \frac{k_B}{2L} \right)^2 \right], \quad (9.54)
$$

where

$$
L_{AB} = L_A + L_B + l_{AB}, \quad (9.55)
$$

$l_{AB}$ is the number of direct links between the nodes of communities A and B, and

$$
k_{AB} = k_A + k_B. \quad (9.56)
$$

After inserting (9.55) and (9.56) into (9.54), we obtain

$$
\Delta M_{AB} = \frac{l_{AB}}{L} - \frac{k_A k_B}{2L^2}, \quad (9.57)
$$

which is (9.13).
The algorithms discussed in this chapter were chosen to illustrate the fundamental ideas and concepts pertaining to community detection. Consequently they are not guaranteed to be neither the fastest nor the most accurate algorithms. Recently two algorithms, called the Louvain algorithm and Infomap have gained popularity, as their accuracy is comparable to the accuracy of the algorithms covered in this chapter but offer better scalability. Consequently we can use them to identify communities in very large networks.

There are many similarities between the two algorithms:

- They both aim to optimize a quality function $Q$. For the Louvain algorithm $Q$ is modularity, $M$, and for Infomap $Q$ is an entropy-based measure called the map equation or $L$.

- Both algorithms use the same optimization procedure.

Given these similarities, we discuss the algorithms together.

THE LOUVAIN ALGORITHM

The $O(N^2)$ computational complexity of the greedy algorithm can be prohibitive for very large networks. A modularity optimization algorithm with better scalability was proposed by Blondel and collaborators [2]. The Louvain algorithm consists of two steps that are repeated iteratively (Figure 9.37):

**Step I**

Start with a weighted network of $N$ nodes, initially assigning each node to a different community. For each node $i$ we evaluate the gain in modularity if we place node $i$ in the community of one of its neighbors $j$. We then move node $i$ in the community for which the modularity gain is the largest, but only if this gain is positive. If no positive gain is found, $i$ stays in its original community. This process is applied to all nodes until no further improvement can be achieved, completing Step I.
The modularity change $\Delta M$ obtained by moving an isolated node $i$ into a community $C$ can be calculated using

$$
\Delta M = \left[ \frac{\sum_{\text{in}} + 2k_{\text{in}}}{2W} - \left( \frac{\sum_{\text{tot}} + k_i}{2W} \right)^2 \right] - \left[ \frac{\sum_{\text{in}}}{2W} - \left( \frac{\sum_{\text{tot}}}{2W} \right)^2 - \left( \frac{k_i}{2W} \right)^2 \right]
$$

(9.58)

where $\sum_{\text{in}}$ is the sum of the weights of the links inside $C$ (which is $L_c$ for an unweighted network); $\sum_{\text{tot}}$ is the sum of the link weights of all nodes in $C$; $k_i$ is the sum of the weights of the links incident to node $i$; $k_{\text{in}}$ is the sum of the weights of the links from $i$ to nodes in $C$ and $W$ is the sum of the weights of all links in the network.

Note that $\Delta M$ is a special case of (9.13), which provides the change in modularity after merging communities A and B. In the current case B is an isolated node. We can use $\Delta M$ to determine the modularity change when $i$ is removed from the community it belonged earlier. For this we calculate $\Delta M$ for merging $i$ with the community $C$ after we excluded $i$ from it. The change after removing $i$ is $-\Delta M$.

### Step II

We construct a new network whose nodes are the communities identified during Step I. The weight of the link between two nodes is the sum of the weight of the links between the nodes in the corresponding communities. Links between nodes of the same community lead to weighted self-loops.

Once Step II is completed, we repeat Steps I - II, calling their combination a pass (Figure 9.37). The number of communities decreases with each pass. The passes are repeated until there are no more changes and maximum modularity is attained.

The main steps of the Louvain algorithm. Each pass consists of two distinct steps:

**Step I**

Modularity is optimized by local changes. We choose a node and calculate the change in modularity, (9.58), if the node joins the community of its immediate neighbors. The figure shows the expected modularity change $\Delta M_{0,i}$ for node 0. Accordingly node 0 will join node 3, as the modularity change for this move is the largest, being $\Delta M_{0,3}=0.032$. This process is repeated for each node, the node colors corresponding to the resulting communities, concluding Step I.

**Step II**

The communities obtained in Step I are aggregated, building a new network of communities. Nodes belonging to the same community are merged into a single node, as shown on the top right. This process will generate self-loops, corresponding to links between nodes in the same community that are now merged into a single node.

The sum of Steps I & II are called a pass. The network obtained after each pass is processed again (Pass 2), until no further increase of modularity is possible. After [2].

Figure 9.37

The Louvain Algorithm
Computational Complexity

The Louvain algorithm is more limited by storage demands than by computational time. The number of computations scale linearly with $L$ for the most time consuming first pass. With subsequent passes over a decreasing number of nodes and links, the complexity of the algorithm is at most $O(L)$. It therefore allows us to identify communities in networks with millions of nodes.

INFOMAP

Introduced by Martin Rosvall and Carl T Bergstrom, Infomap exploits data compression for community identification (Figure 9.38) [44-46]. It does it by optimizing a quality function for community detection in directed and weighted networks, called the map equation.

Consider a network partitioned into $n_c$ communities. We wish to encode in the most efficient fashion the trajectory of a random walker on this network. In other words, we want to describe the trajectory with the smallest number of symbols. The ideal code should take advantage of the fact that the random walker tends to get trapped into communities, staying there for a long time (Figure 9.38c).

To achieve this coding we assign:

- One code to each community (index codebook). For example the purple community in Figure 9.38c is assigned the code 111.
- Codewords for each node within each community. For example the top left node in (c) is assigned 001. Note that the same node code can be reused in different communities.
- Exit codes that mark when the walker leaves a community, like 0001 for the purple community in (c).

The goal, therefore, is to build a code that offers the shortest description of the random walk. Once we have this code, we can identify the network’s community structure by reading the index codebook, which is uniquely as-

Figure 9.38

From Data Compression to Communities

Infomap detect communities by compressing the movement of a random walker on a network.

(a) The orange line shows the trajectory of a random walker on a small network. We want to describe this trajectory with a minimal number of symbols, which we can achieve by assigning repeatedly used structures (communities) short and unique names.

(b) We start by giving a unique name to each node. This is derived using a Huffman coding, a data compression algorithm that assigns each node a code using the estimated probability that the random walk visits that node. The 314 bits under the network describe the sample trajectory of the random walker shown in (a), starting with 1111100 for the first node of the walk in the upper left corner, 1100 for the second node, etc., and ending with 00011 for the last node on the walk in the lower right corner.

(c) The figure shows a two-level encoding of the random walk, in which each community receives a unique name, but the name of nodes within communities are reused. This code yields on average a 32% shorter coding. The codes naming the communities and the codes used to indicate an exit from each community are shown to the left and the right of the arrows under the network, respectively. Using this code, we can describe the walk in (a) by the 243 bits shown under the network in (c). The first three bits 111 indicate that the walk begins in the red community, the code 0000 specifies the first node of the walk, etc.

(d) By reporting only the community names, and not the locations of each node within the communities, we obtain an efficient coarse graining of the network, which corresponds to its community structure.
signed to each community (Figure 9.38c).

The optimal code is obtained by finding the minimum of the map equation

$$\mathcal{L} = qH(Q) + \sum_{c=1}^{n_c} p_c H(P_c)$$  \hspace{1cm} (9.59)

In a nutshell, the first term of (9.59) gives the average number of bits necessary to describe the movement between communities where $q$ is the probability that the random walker switches communities during a given step.

The second term gives the average number of bits necessary to describe movement within communities. Here $H(P_c)$ is the entropy of within-community movements — including an “exit code” to capture the departure from a community $i$.

The specific terms of the maps equation and their calculation in terms of the probabilities capturing the movement of a random walker on a network, is somewhat involved. They are described in detail in Ref [44-46]. Online Resource 9.3 offers an interactive tool to illustrate the mechanism behind (9.59) and its use.

At the end $\mathcal{L}$ serves as a quality function, that takes up a specific value for a particular partition of the network into communities. To find the best partition, we must minimize $\mathcal{L}$ over all possible partitions. The popular implementation of this optimization procedure follows Steps I and II of the Louvain algorithm: We assign each node to a separate community, and we systematically join neighboring nodes into modules if the move decreases $\mathcal{L}$. After each move $\mathcal{L}$ is updated using (9.59). The obtained communities are joined into supercommunities, finishing a pass, after which the algorithm is restarted on the new, reduced network.

**Computational Complexity**

The computational complexity of Infomap is determined by the procedure used to minimize the map equation $\mathcal{L}$. If we use the Louvain procedure, the computational complexity is the same as that of the Louvain algorithm, i.e. at most $O(L\log L)$ or $O(N\log N)$ for a sparse graph.

In summary, the Louvain algorithm and Infomap offer tools for fast community identification. Their accuracy across benchmarks is comparable to the accuracy of the algorithms discussed throughout this chapter (Figure 9.28).
In this section we derive the percolation threshold \((9.20)\) for clique percolation on a random network and discuss the main steps of the CFinder algorithm (Figure 9.39).

When we roll a \(k\)-clique to an adjacent \(k\)-clique by relocating one of its nodes, the expectation value of the number of adjacent \(k\)-cliques for the template to roll further should equal exactly one at the percolation threshold (Figure 9.20). Indeed, a smaller than one expectation value will result in a premature end of the \(k\)-clique percolation clusters, because starting from any \(k\)-clique, the rolling would quickly come to a halt. Consequently the size of the clusters would decay exponentially. A larger than one expectation value, on the other hand, allows the clique community to grow indefinitely, guaranteeing that we have a giant cluster in the system.

The above expectation value is provided by

\[
(k - 1)(N - k - 1)^{k-1},
\]

where the term \((k-1)\) counts the number of nodes of the template that can be selected for the next relocation; the term \((N-k-1)^{k-1}\) counts the number of potential destinations for this relocation, out of which only the fraction \(p^{k-1}\) is acceptable, because each of the new \(k-1\) edges (associated with the relocation) must exist in order to obtain a new \(k\)-clique. For large \(N\), \((9.63)\) simplifies to

\[
(k-1)Np^{k-1} = 1,
\]

which leads to \((9.16)\).
Figure 9.39
CFinder algorithm

The main steps of the CFinder algorithm.

(a) Starting from the network shown in the figure, our goal is to identify all cliques. All five $k=3$ cliques present in the network are highlighted.

(b) The overlap matrix $O$ of the $k=3$ cliques. This matrix is viewed as an adjacency matrix of a network whose nodes are the cliques of the original network. The matrix indicates that we have two connected components, one consisting of cliques $(1,2)$ and the other of cliques $(3,4,5)$. The connected components of this network map into the communities of the original network.

(c) The two clique communities predicted by the adjacency matrix.

(d) The two clique communities shown in (c), mapped on the original network.


