ALBERT-LÁSZLÓ BARABÁSI

NETWORK SCIENCE THE SCALE-FREE PROPERTY



ACKNOWLEDGEMENTS

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works.





chitect, he deploys insights from engineering, physics, chemistry, aeronautics, and materials science, using networks as a source of inspiration and metaphor. The image shows his work displayed in the Miami Art Museum, an example of the artist's take on complex net-

INTRODUCTION

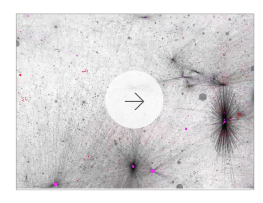
The World Wide Web is a network whose nodes are documents and the links are the uniform resource locators (URLs) that allow us to "surf" with a click from one web document to the other. With an estimated size of over one trillion documents ($N \approx 10^{12}$), the Web is the largest network humanity has ever built. It exceeds in size even the human brain ($N \approx 10^{11}$ neurons).

It is difficult to overstate the importance of the World Wide Web in our daily life. Similarly, we cannot exaggerate the role the WWW played in the development of network theory: it facilitated the discovery of a number of fundamental network characteristics and became a standard testbed for most network measures.

We can use a software called a *crawler* to map out the Web's wiring diagram. A crawler can start from any web document, identifying the links (URLs) on it. Next it downloads the documents these links point to and identifies the links on these documents, and so on. This process iteratively returns a local map of the Web. Search engines like Google or Bing operate crawlers to find and index new documents and to maintain a detailed map of the WWW.

The first map of the WWW obtained with the explicit goal of understanding the structure of the network behind it was generated by Hawoong Jeong at University of Notre Dame. He mapped out the nd.edu domain [1], consisting of about 300,000 documents and 1.5 million links (Online Resource 4.1). The purpose of the map was to compare the properties of the Web graph to the random network model. Indeed, in 1998 there were reasons to believe that the WWW could be well approximated by a random network. The content of each document reflects the personal and professional interests of its creator, from individuals to organizations. Given the diversity of these interests, the links on these documents might appear to point to randomly chosen documents.

A quick look at the map in Figure 4.1 supports this view: There appears to be considerable randomness behind the Web's wiring diagram. Yet, a



Online Resource 4.1

Zooming into the World Wide Web

Watch an online video that zooms into the WWW sample that has lead to the discovery of the scale-free property [1]. This is the network featured in Table 2.1 and shown in Figure 4.1, whose characteristics are tested throughout this book.



closer inspection reveals some puzzling differences between this map and a random network. Indeed, in a random network highly connected nodes, or hubs, are effectively forbidden. In contrast in Figure 4.1 numerous small-degree nodes coexist with a few hubs, nodes with an exceptionally large number of links.

In this chapter we show that hubs are not unique to the Web, but we encounter them in most real networks. They represent a signature of a deeper organizing principle that we call the scale-free property. We therefore explore the degree distribution of real networks, which allows us to uncover and characterize scale-free network. The analytical and empirical results discussed here represent the foundations of the modeling efforts the rest of this book is based on. Indeed, we will come to see that no matter what network property we are interested in, from communities to spreading processes, it must be inspected in the light of the network's degree distribution.

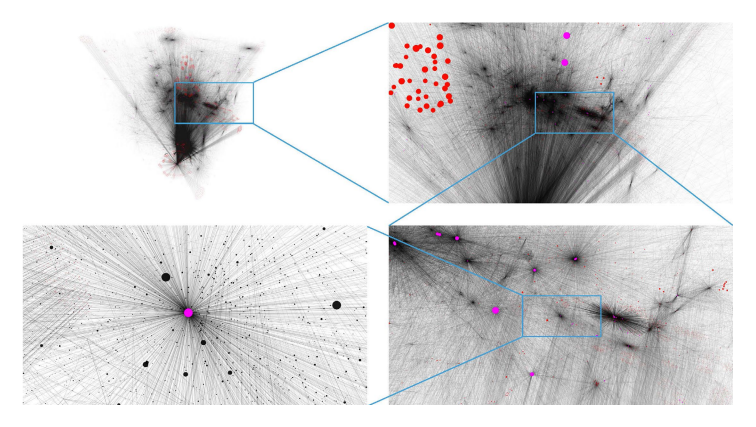


Figure 4.1
The Topology of the World Wide Web

Snapshots of the World Wide Web sample mapped out by Hawoong Jeong in 1998 [1]. The sequence of images show an increasingly magnified local region of the network. The first panel displays all 325,729 nodes, offering a global view of the full dataset. Nodes with more than 50 links are shown in red and nodes with more than 500 links in purple. The closeups reveal the presence of a few highly connected nodes, called *hubs*, that accompany scale-free networks. Courtesy of M. Martino.

POWER LAWS AND SCALE-FREE NETWORKS

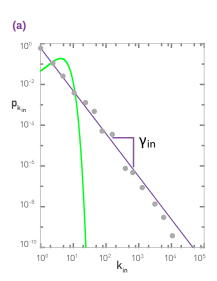
If the WWW were to be a random network, the degrees of the Web documents should follow a Poisson distribution. Yet, as Figure 4.2 indicates, the Poisson form offers a poor fit for the WWW's degree distribution. Instead on a log-log scale the data points form an approximate straight line, suggesting that the degree distribution of the WWW is well approximated with

$$p_{k} \sim k^{-\gamma}. \tag{4.1}$$

Equation (4.1) is called a *power law distribution* and the exponent γ is its *degree exponent* (BOX 4.1). If we take a logarithm of (4.1), we obtain

$$\log p_k \sim -\gamma \log k. \tag{4.2}$$

If (4.1) holds, $\log p_k$ is expected to depend linearly on $\log k$, the slope of this line being the degree exponent γ (Figure 4.2).



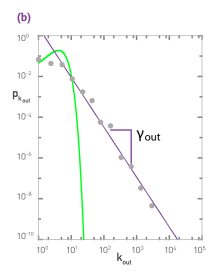


Figure 4.2
The Degree Distribution of the WWW

The incoming **(a)** and outgoing **(b)** degree distribution of the WWW sample mapped in the 1999 study of Albert *et al.* [1]. The degree distribution is shown on double logarithmic axis (log-log plot), in which a power law follows a straight line. The symbols correspond to the empirical data and the line corresponds to the power-law fit, with degree exponents γ_{in} = 2.1 and γ_{out} = 2.45. We also show as a green line the degree distribution predicted by a Poisson function with the average degree $\langle k_{in} \rangle = \langle k_{out} \rangle = 4.60$ of the WWW sample.

The WWW is a directed network, hence each document is characterized by an out-degree k_{out} , representing the number of links that point from the document to other documents, and an in-degree k_{in} , representing the number of other documents that point to the selected document. We must therefore distinguish two degree distributions: the probability that a randomly chosen document points to $k_{\scriptscriptstyle out}$ web documents, or $p_{\scriptscriptstyle k_{\scriptscriptstyle out}}$, and the probability that a randomly chosen node has k_{in} web documents pointing to it, or $p_{\mathbf{k}_{in}}$. In the case of the WWW both $p_{\mathbf{k}_{in}}$ and $p_{\mathbf{k}_{out}}$ can be approximated by a power law

$$p_{k_{-}} \sim k^{-\gamma_{in}}$$
 (4.3)

$$p_{k_{out}} \sim k^{-\gamma_{out}}$$
, (4.4)

where γ_{in} and γ_{out} are the degree exponents for the in- and out-degrees, respectively (Figure 4.2). In general γ_{in} can differ from γ_{out} . For example, in Figure 4.1 we have $\gamma_{in} \approx 2.1$ and $\gamma_{out} \approx 2.45$.

The empirical results shown in Figure 4.2 document the existence of a network whose degree distribution is quite different from the Poisson distribution characterizing random networks. We will call such networks scale-free, defined as [2]:

A scale-free network is a network whose degree distribution follows a power law.

As Figure 4.2 indicates, for the WWW the power law persists for almost four orders of magnitude, prompting us to call the Web graph scale-free network. In this case the scale-free property applies to both in and out-degrees.

To better understand the scale-free property, we have to define the power-law distribution in more precise terms. Therefore next we discuss the discrete and the continuum formalisms used throughout this book.

Discrete Formalism

As node degrees are positive integers, k = 0, 1, 2, ..., the discrete formalism provides the probability p_k that a node has exactly k links

$$p_{k} = Ck^{-\gamma}. \tag{4.5}$$

The constant C is determined by the normalization condition

$$\sum_{k=1}^{\infty} p_k = 1.$$

$$C \sum_{k=1}^{\infty} k^{-\gamma} = 1,$$
(4.6)

Using (4.5) we obtain,

$$C\sum_{i=1}^{\infty}k^{-\gamma}=I,$$

hence

$$C = \frac{1}{\sum_{k=1}^{\infty} k^{-\gamma}} = \frac{1}{\zeta(\gamma)},\tag{4.7}$$

where $\zeta(\gamma)$ is the Riemann-zeta function. Thus for k>0 the discrete power-law distribution has the form

$$\rho_{k} = \frac{k^{-\gamma}}{\zeta(\gamma)}. (4.8)$$

Note that (4.8) diverges at k=0. If needed, we can separately specify p_o , representing the fraction of nodes that have no links to other nodes. In that case the calculation of C in (4.7) needs to incorporate p_o .

Continuum Formalism

In analytical calculations it is often convenient to assume that the degrees can have any positive real value. In this case we write the power-law degree distribution as

$$p(k) = Ck^{-\gamma}. (4.9)$$

Using the normalization condition

$$\int_{k_{\min}}^{\infty} p(k)dk = I \tag{4.10}$$

we obtain

$$C = \frac{1}{\int_{k_{\min}}^{\infty} k^{-\gamma} dk} = (\gamma - 1)k_{\min}^{\gamma - 1}.$$
 (4.11)

Therefore in the continuum formalism the degree distribution has the form

$$p(k) = (\gamma - 1)k_{min}^{\gamma - 1} k^{-\gamma}. \tag{4.12}$$

Here k_{min} is the smallest degree for which the power law (4.8) holds.

Note that p_k encountered in the discrete formalism has a precise meaning: it is the probability that a randomly selected node has degree k. In contrast, only the integral of p(k) encountered in the continuum formalism has a physical interpretation:

$$\int_{k}^{k_2} p(k) dk \tag{4.13}$$

is the probability that a randomly chosen node has degree between $k_{\!\scriptscriptstyle I}$ and $k_{\!\scriptscriptstyle 2}\!.$

In summary, networks whose degree distribution follows a power law are called scale-free networks. If a network is directed, the scale-free property applies separately to the in- and the out-degrees. To mathematically study the properties of scale-free networks, we can use either the discrete or the continuum formalism. The scale-free property is independent of the formalism we use.

BOX 4.1

THE 80/20 RULE AND THE TOP ONE PERCENT

Vilfredo Pareto, a 19th century economist, noticed that in Italy a few wealthy individuals earned most of the money, while the majority of the population earned rather small amounts. He connected this disparity to the observation that incomes follow a power law, representing the first known report of a power-law distribution [3]. His finding entered the popular literature as the 80/20 rule: Roughly 80 percent of money is earned by only 20 percent of the population.

The 80/20 rule emerges in many areas. For example in management it is often stated that 80 percent of profits are produced by only 20 percent of the employees. Similarly, 80 percent of decisions are made during 20 percent of meeting time.

The 80/20 rule is present in networks as well: 80 percent of links on the Web point to only 15 percent of webpages; 80 percent of citations go to only 38 percent of scientists; 80 percent of links in Hollywood are connected to 30 percent of actors [4]. Most quantities following a power law distribution obey the 80/20 rule.

During the 2009 economic crisis power laws gained a new meaning: The Occupy Wall Street Movement draw attention to the fact that in the US 1% of the population earns a disproportionate 15% of the total US income. This 1% phenomena, a signature of a profound income disparity, is again a consequence of the power-law nature of the income distribution.



Figure 4.3 Vilfredo Federico Damaso Pareto (1848 – 1923)

Italian economist, political scientist, and philosopher, who had important contributions to our understanding of income distribution and to the analysis of individual choices. A number of fundamental principles are named after him, like *Pareto efficiency*, *Pareto distribution* (another name for a power-law distribution), the *Pareto principle* (or 80/20 law).

HUBS

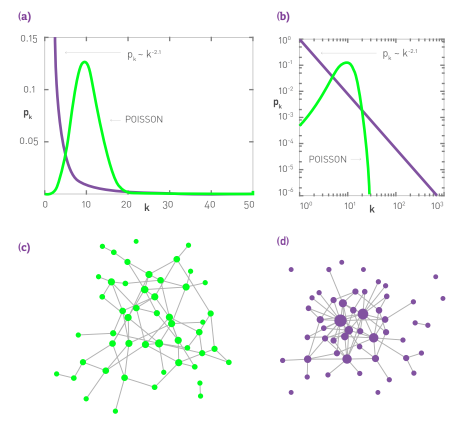
The main difference between a random and a scale-free network comes in the tail of the degree distribution, representing the high-k region of p_k . To illustrate this, in Figure 4.4 we compare a power law with a Poisson function. We find that:

- For small *k* the power law is above the Poisson function, indicating that a scale-free network has a large number of small degree nodes, most of which are absent in a random network.
- For k in the vicinity of $\langle k \rangle$ the Poisson distribution is above the power law, indicating that in a random network there is an excess of nodes with degree $k \approx \langle k \rangle$.
- For large k the power law is again above the Poisson curve. The difference is particularly visible if we show p_k on a log-log plot (Figure 4.4b), indicating that the probability of observing a high-degree node, or hub, is several orders of magnitude higher in a scale-free than in a random network.

Let us use the WWW to illustrate the magnitude of these differences. The probability to have a node with k=100 is about $p_{100} \approx 10^{-94}$ in a Poisson distribution while it is about $p_{100} \approx 4 \times 10^{-4}$ if p_k follows a power law. Consequently, if the WWW were to be a random network with < k > = 4.6 and size $N \approx 10^{12}$, we would expect

$$N_{k \ge 100} = 10^{12} \sum_{k=100}^{\infty} \frac{(4.6)^k}{k!} e^{-4.6} \simeq 10^{-82}$$
 (4.14)

nodes with at least 100 links, or effectively none. In contrast, given the WWW's power law degree distribution, with $\gamma_{in} = 2.1$ we have $N_{k \ge 100} = 4 \times 10^9$, i.e. more than four billion nodes with degree $k \ge 100$.



The Largest Hub

All real networks are finite. The size of the WWW is estimated to be $N \approx 10^{12}$ nodes; the size of the social network is the Earth's population, about $N \approx 7 \times 10^9$. These numbers are huge, but finite. Other networks pale in comparison: The genetic network in a human cell has approximately 20,000 genes while the metabolic network of the *E. Coli* bacteria has only about a thousand metabolites. This prompts us to ask: How does the network size affect the size of its hubs? To answer this we calculate the maximum degree, k_{max} , called the *natural cutoff* of the degree distribution p_k . It represents the expected size of the largest hub in a network.

It is instructive to perform the calculation first for the exponential distribution

$$p(k) = Ce^{-\lambda k}$$
.

For a network with minimum degree $k_{\mbox{\tiny min}}$ the normalization condition

$$\int_{k_{min}}^{\infty} p(k)dk = 1 \tag{4.15}$$

provides $C = \lambda e^{\lambda k_{\min}}$. To calculate k_{max} we assume that in a network of N nodes we expect at most one node in the (k_{max}, ∞) regime (ADVANCED TOPICS 3.B). In other words the probability to observe a node whose degree exceeds k_{max} is 1/N:

$$\int_{k_{max}}^{\infty} p(k)dk = \frac{1}{N}.$$
(4.16)

Figure 4.4
Poisson vs. Power-law Distributions

- (a) Comparing a Poisson function with a power-law function (γ = 2.1) on a linear plot. Both distributions have $\langle k \rangle$ = 11.
- (b) The same curves as in (a), but shown on a log-log plot, allowing us to inspect the difference between the two functions in the high-k regime.
- (c) A random network with $\langle k \rangle = 3$ and N = 50, illustrating that most nodes have comparable degree $k \approx \langle k \rangle$.
- (d) A scale-free network with γ =2.1 and $\langle k \rangle$ =3, illustrating that numerous small-degree nodes coexist with a few highly connected hubs. The size of each node is proportional to its degree.

Equation (4.16) yields

$$k_{\text{max}} = k_{\text{min}} + \frac{\ln N}{\lambda}.\tag{4.17}$$

As $\ln N$ is a slow function of the system size, (4.17) tells us that the maximum degree will not be significantly different from k_{min} . For a Poisson degree distribution the calculation is a bit more involved, but the obtained dependence of k_{max} on N is even slower than the logarithmic dependence predicted by (4.17) (ADVANCED TOPICS 3.B).

For a scale-free network, according to (4.12) and (4.16), the natural cutoff follows

$$k_{max} = k_{min} N^{\frac{1}{\gamma - 1}}.$$
 (4.18)

Hence the larger a network, the larger is the degree of its biggest hub. The polynomial dependence of k_{max} on N implies that in a large scale-free network there can be orders of magnitude differences in size between the smallest node, k_{min} , and the biggest hub, k_{max} (Figure 4.5).

To illustrate the difference in the maximum degree of an exponential and a scale-free network let us return to the WWW sample of Figure 4.1, consisting of $N \approx 3 \times 10^5$ nodes. As $k_{min} = 1$, if the degree distribution were to follow an exponential, (4.17) predicts that the maximum degree should be $k_{max} \approx 14$ for $\lambda=1$. In a scale-free network of similar size and $\gamma=2.1$, (4.18) predicts $k_{max} \approx 95,000$, a remarkable difference. Note that the largest in-degree of the WWW map of Figure 4.1 is 10,721, which is comparable to k_{max} predicted by a scale-free network. This reinforces our conclusion that in a random network hubs are effectively forbidden, while in scale-free networks they are naturally present.

In summary the key difference between a random and a scale-free network is rooted in the different shape of the Poisson and of the power-law function: In a random network most nodes have comparable degrees and hence hubs are forbidden. Hubs are not only tolerated, but are expected in scale-free networks (Figure 4.6). Furthermore, the more nodes a scale-free network has, the larger are its hubs. Indeed, the size of the hubs grows polynomially with network size, hence they can grow quite large in scale-free networks. In contrast in a random network the size of the largest node grows logarithmically or slower with N, implying that hubs will be tiny even in a very large random network.

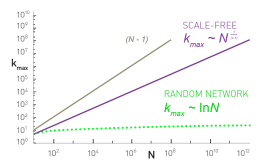
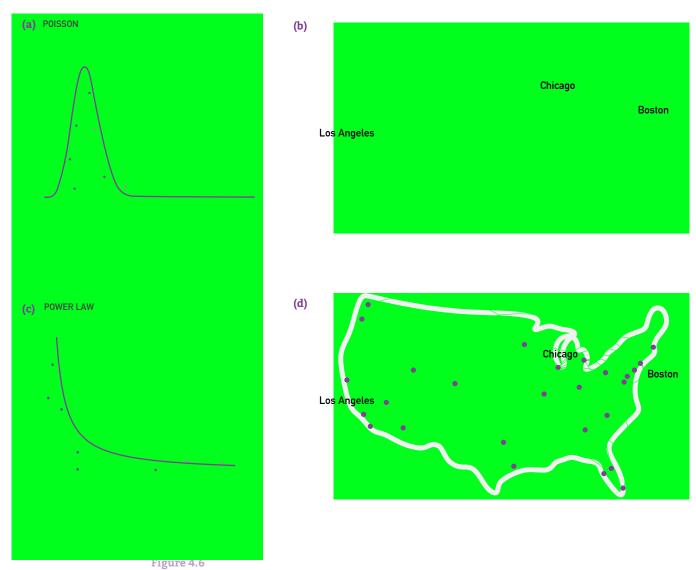


Figure 4.5 Hubs are Large in Scale-free Networks

The estimated degree of the largest node (natural cutoff) in scale-free and random networks with the same average degree $\langle k \rangle = 3$. For the scale-free network we chose $\gamma = 2.5$. For comparison, we also show the linear behavior, $k_{max} \sim N-1$, expected for a complete network. Overall, hubs in a scale-free network are several orders of magnitude larger than the biggest node in a random network with the same N and $\langle k \rangle$.



Random vs. Scale-free Networks

- (a) The degrees of a random network follow a Poisson distribution, rather similar to a bell curve. Therefore most nodes have comparable degrees and nodes with a large number of links are absent.
- **(b)** A random network looks a bit like the national highway network in which nodes are cities and links are the major highways. There are no cities with hundreds of highways and no city is disconnected from the highway system.
- (c) In a network with a power-law degree distribution most nodes have only a few links. These numerous small nodes are held together by a few highly connected hubs.
- (d) A scale-free network looks like the air-traffic network, whose nodes are airports and links are the direct flights between them. Most airports are tiny, with only a few flights. Yet, we have a few very large airports, like Chicago or Los Angeles, that act as major hubs, connecting many smaller airports.

Once hubs are present, they change the way we navigate the network. For example, if we travel from Boston to Los Angeles by car, we must drive through many cities. On the airplane network, however, we can reach most destinations via a single hub, like Chicago. After [4].

THE MEANING OF SCALE-FREE

The term "scale-free" is rooted in a branch of statistical physics called the *theory of phase transitions* that extensively explored power laws in the 1960s and 1970s (ADVANCED TOPICS 3.F). To best understand the meaning of the scale-free term, we need to familiarize ourselves with the moments of the degree distribution.

The n^{th} moment of the degree distribution is defined as

$$\langle k^n \rangle = \sum_{k_{\min}}^{\infty} k^n p_k \approx \int_{k_{\min}}^{\infty} k^n p(k) dk.$$
 (4.19)

The lower moments have important interpretation:

- n=1: The first moment is the average degree, $\langle k \rangle$.
- n=2: The second moment, $\langle k^2 \rangle$, helps us calculate the *variance* $\sigma^2 = \langle k^2 \rangle$ $\langle k \rangle^2$, measuring the spread in the degrees. Its square root, σ , is the *standard deviation*.
- n=3: The third moment, $\langle k^3 \rangle$, determines the *skewness* of a distribution, telling us how symmetric is p_k around the average $\langle k \rangle$.

For a scale-free network the n^{th} moment of the degree distribution is

$$\langle k^n \rangle = \int_{k_{\min}}^{k_{\max}} k^n p(k) dk = C \frac{k_{\max}^{n-\gamma+1} - k_{\min}^{n-\gamma+1}}{n-\gamma+1}.$$
 (4.20)

While typically k_{\min} is fixed, the degree of the largest hub, k_{\max} , increases with the system size, following (4.18). Hence to understand the behavior of $\langle k^n \rangle$ we need to take the asymptotic limit $k_{\max} \to \infty$ in (4.20), probing the properties of very large networks. In this limit (4.20) predicts that the value of $\langle k^n \rangle$ depends on the interplay between n and γ :

- If $n-\gamma+1\leq 0$ then the first term on the r.h.s. of (4.20), $k_{\max}^{n-\gamma+1}$, goes to zero as k_{\max} increases. Therefore all moments that satisfy $n\leq \gamma-1$ are finite.
- If $n-\gamma+1>0$ then $\langle k^n\rangle$ goes to infinity as $k_{\max}\to\infty$. Therefore all mo-

ments larger than γ-1 diverge.

For many scale-free networks the degree exponent γ is between 2 and 3 (Table 4.1). Hence for these in the $N \to \infty$ limit the first moment $\langle k \rangle$ is finite, but the second and higher moments, $\langle k^2 \rangle$, $\langle k^3 \rangle$, go to infinity. This divergence helps us understand the origin of the "scale-free" term. Indeed, if the degrees follow a normal distribution, then the degree of a randomly chosen node is typically in the range

$$k = \langle k \rangle \pm \sigma_k \,. \tag{4.21}$$

Yet, the average degree < k > and the standard deviation σ_k have rather different magnitude in random and in scale-free networks:

• Random Networks Have a Scale

For a random network with a Poisson degree distribution $\sigma_k = \langle k \rangle^{1/2}$, which is always smaller than $\langle k \rangle$. Hence the network's nodes have degrees in the range $k = \langle k \rangle \pm \langle k \rangle^{1/2}$. In other words nodes in a random network have comparable degrees and the average degree $\langle k \rangle$ serves as the "scale" of a random network.

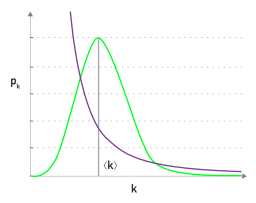
• Scale-free Networks Lack a Scale

For a network with a power-law degree distribution with $\gamma < 3$ the first moment is finite but the second moment is infinite. The divergence of $\langle k^2 \rangle$ (and of σ_k) for large N indicates that the fluctuations around the average can be arbitrary large. This means that when we randomly choose a node, we do not know what to expect: The selected node's degree could be tiny or arbitrarily large. Hence networks with $\gamma < 3$ do not have a meaningful internal scale, but are "scale-free" (Figure 4.7).

For example the average degree of the WWW sample is $\langle k \rangle$ = 4.60 (Table 4.1). Given that $\gamma \approx 2.1$, the second moment diverges, which means that our expectation for the in-degree of a randomly chosen WWW document is k=4.60 $\pm \infty$ in the $N \to \infty$ limit. That is, a randomly chosen web document could easily yield a document of degree one or two, as 74.02% of nodes have in-degree less than $\langle k \rangle$. Yet, it could also yield a node with hundreds of millions of links, like google.com or facebook. com.

Strictly speaking $\langle k^2 \rangle$ diverges only in the $N \to \infty$ limit. Yet, the divergence is relevant for finite networks as well. To illustrate this, Table 4.1 lists $\langle k^2 \rangle$ and Figure 4.8 shows the standard deviation σ for ten real networks. For most of these networks σ is significantly larger than $\langle k \rangle$, documenting large variations in node degrees. For example, the degree of a randomly chosen node in the WWW sample is k_{in} = 4.60 ± 1546, indicating once again that the average is not informative.

In summary, the scale-free name captures the lack of an internal scale, a consequence of the fact that nodes with widely different degrees coexist in the same network. This feature distinguishes scale-free networks from lattices, in which all nodes have exactly the same degree ($\sigma=0$), or from random networks, whose degrees vary in a narrow range ($\sigma=\langle k\rangle^{1/2}$). As we



Random Network Randomly chosen node: $k = \langle k \rangle \pm \langle k \rangle^{1/2}$ Scale: $\langle k \rangle$

Scale-Free Network Randomly chosen node: $k = \left\langle k \right\rangle \pm \infty$ Scale: none

Figure 4.7 Lack of an Internal Scale

For any exponentially bounded distribution, like a Poisson or a Gaussian, the degree of a randomly chosen node is in the vicinity of $\langle k \rangle$. Hence $\langle k \rangle$ serves as the network's scale. For a power law distribution the second moment can diverge, and the degree of a randomly chosen node can be significantly different from $\langle k \rangle$. Hence $\langle k \rangle$ does not serve as an intrinsic scale. As a network with a power law degree distribution lacks an intrinsic scale, we

will see in the coming chapters, this divergence is the origin of some of the most intriguing properties of scale-free networks, from their robustness to random failures to the anomalous spread of viruses.

NETWORK	N	L	$\langle k \rangle$	$\langle k_{in}^2 \rangle$	$\langle k_{out}^2 \rangle$	$\langle k^2 \rangle$	$\gamma_{\it in}$	γ_{out}	γ
Internet	192,244	609,066	6.34	-	-	240.1	-	_	3.42*
www	325,729	1,497,134	4.60	1546.0	482.4	-	2.00	2.31	-
Power Grid	4,941	6,594	2.67	-	-	10.3	-	-	Exp.
Mobile Phone Calls	36,595	91,826	2.51	12.0	11.7	-	4.69*	5.01*	-
Email	57,194	103,731	1.81	94.7	1163.9	-	3.43*	2.03*	-
Science Collaboration	23,133	93,439	8.08	-	-	178.2	-	-	3.35*
Actor Network	702,388	29,397,908	83.71	-	-	47,353.7	-	-	2.12*
Citation Network	449,673	4,689,479	10.43	971.5	198.8	-	3.03**	4.00*	-
E. Coli Metabolism	1,039	5,802	5.58	535.7	396.7	-	2.43*	2.9 0*	_
Protein Interactions	2,018	2,930	2.90	-	-	32.3	-	-	2.89*

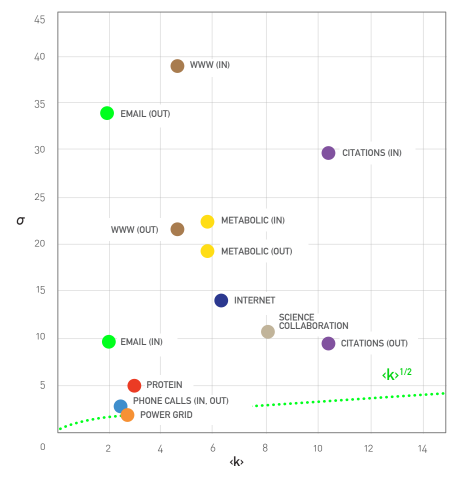


Table 4.1

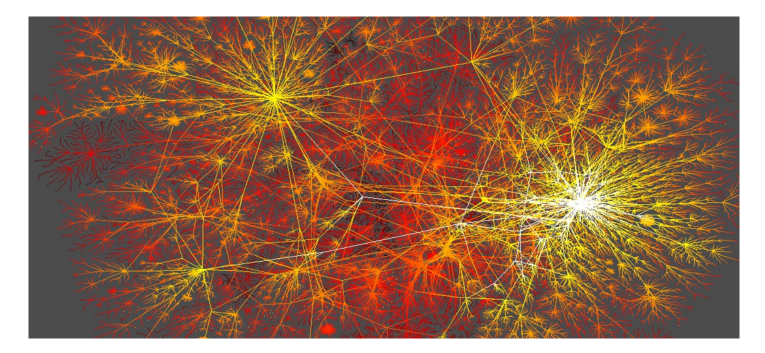
Degree Fluctuations in Real Networks

The table shows the first $\langle k \rangle$ and the second moment $\langle k^2 \rangle$ ($\langle k_{in}^2 \rangle$ and $\langle k_{out}^2 \rangle$ for directed networks) for ten reference networks. For directed networks we list $\langle k \rangle = \langle k_{in} \rangle = \langle k_{out} \rangle$. We also list the estimated degree exponent, y, for each network, determined using the procedure discussed in ADVANCED TOPICS 4.A. The stars next to the reported values indicate the confidence of the fit to the degree distribution. That is, ' means that the fit shows statistical confidence for a power-law $(k^{-\gamma})$; while ** marks statistical confidence for a fit (4.39) with an exponential cutoff. Note that the power grid is not scalefree. For this network a degree distribution of the form $e^{-\lambda k}$ offers a statistically significant fit, which is why we placed an "Exp" in the last column.

Figure 4.8
Standard Deviation is Large in Real Networks

For a random network the standard deviation follows $\sigma = \langle k \rangle^{1/2}$ shown as a green dashed line on the figure. The symbols show σ for nine of the ten reference networks, calculated using the values shown in Table 4.1. The actor network has a very large $\langle k \rangle$ and σ , hence it omitted for clarity. For each network σ is larger than the value expected for a random network with the same $\langle k \rangle$. The only exception is the power grid, which is not scale-free. While the phone call network is scale-free, it has a large γ , hence it is well approximated by a random network.

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While the terms WWW and Internet are often used interchangeably in the media, they refer to different systems. The WWW is an information network, whose nodes are documents and links are URLs. In contrast the Internet is an infrastructural network, whose nodes are computers called routers and whose links correspond to physical connections, like copper and optical cables or wireless links.

This difference has important consequences: The cost of linking a Boston-based web page to a document residing on the same computer or to one on a Budapest-based computer is the same. In contrast, establishing a direct Internet link between routers in Boston and Budapest would require us to lay a cable between North America and Europe, which is prohibitively expensive. Despite these differences, the degree distribution of both networks is well approximated by a power law [1, 5, 6]. The signatures of the Internet's scale-free nature are visible in Figure 4.9, showing that a

Figure 4.9
The topology of the Internet

An iconic representation of the Internet topology at the beginning of the 21st century. The image was produced by CAIDA, an organization based at University of California in San Diego, devoted to collect, analyze, and visualize Internet data. The map illustrates the Internet's scale-free nature: A few highly connected hubs hold together numerous small nodes.

few high-degree routers hold together a large number of routers with only a few links.

In the past decade many real networks of major scientific, technological and societal importance were found to display the scale-free property. This is illustrated in Figure 4.10, where we show the degree distribution of an infrastructural network (Internet), a biological network (protein interactions), a communication network (emails) and a network characterizing scientific communications (citations). For each network the degree distribution significantly deviates from a Poisson distribution, being better approximated with a power law.

The diversity of the systems that share the scale-free property is remarkable (BOX 4.2). Indeed, the WWW is a man-made network with a history of little more than two decades, while the protein interaction network is the product of four billion years of evolution. In some of these networks the nodes are molecules, in others they are computers. It is this diversity that prompts us to call the scale-free property a *universal* network characteristic.

From the perspective of a researcher, a crucial question is the following: How do we know if a network is scale-free? On one end, a quick look at the degree distribution will immediately reveal whether the network could be scale-free: In scale-free networks the degrees of the smallest and the largest nodes are widely different, often spanning several orders of magnitude. In contrast, these nodes have comparable degrees in a random network. As the value of the degree exponent plays an important role in predicting various network properties, we need tools to fit the \boldsymbol{p}_k distribution and to estimate γ . This prompts us to address several issues pertaining to plotting and fitting power laws:

Plotting the Degree Distribution

The degree distributions shown in this chapter are plotted on a double logarithmic scale, often called a log-log plot. The main reason is that when we have nodes with widely different degrees, a linear plot is unable to display them all. To obtain the clean-looking degree distributions shown throughout this book we use logarithmic binning, ensuring that each datapoint has sufficient number of observations behind it. The practical tips for plotting a network's degree distribution are discussed in ADVANCED TOPICS 4.B.

Measuring the Degree Exponent

A quick estimate of the degree exponent can be obtained by fitting a straight line to p_k on a log-log plot. Yet, this approach can be affected by systematic biases, resulting in an incorrect γ . The statistical tools available to estimate γ are discussed in ADVANCED TOPICS 4.C.

The Shape of p, for Real Networks

Many degree distributions observed in real networks deviate from a pure power law. These deviations can be attributed to data incompleteness or data collection biases, but can also carry important information about processes that contribute to the emergence of a particular network. In ADVANCED TOPICS 4.B we discuss some of these deviations and in CHAPTER 6 we explore their origins.

In summary, since the 1999 discovery of the scale-free nature of the WWW, a large number of real networks of scientific and technological interest have been found to be scale-free, from biological to social and linguistic networks (BOX 4.2). This does not mean that all networks are scale-free. Indeed, many important networks, from the power grid to networks observed in materials science, do not display the scale-free property (BOX 4.3).

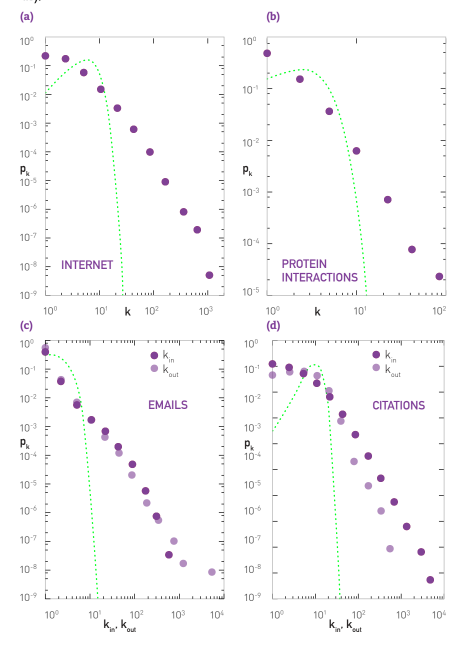


Figure 4.10
Many Real Networks are Scale-free

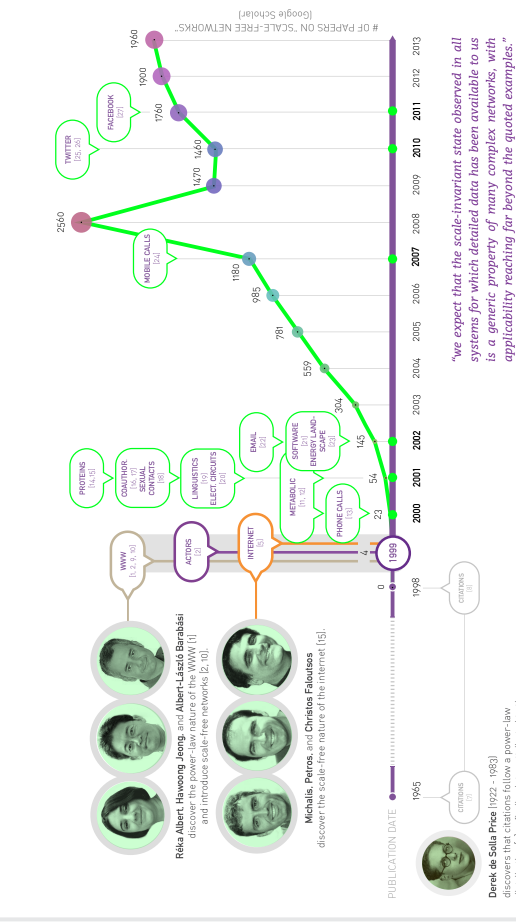
The degree distribution of four networks listed in Table 4.1.

- (a) Internet at the router level.
- (b) Protein-protein interaction network.
- (c) Email network.
- (d) Citation network.

In each panel the green dotted line shows the Poisson distribution with the same $\langle k \rangle$ as the real network, illustrating that the random network model cannot account for the observed $p_k.$ For directed networks we show separately the incoming and outgoing degree distributions.

B0X 4.2

TIMELINE: SCALE-FREE NETWORKS



Barabási and Albert, 1999

distribution [7], a finding later attributed to the scale-free nature of the citation network [2].

BOX 4.3

NOT ALL NETWORK ARE SCALE-FREE

The ubiquity of the scale-free property does not mean that *all* real networks are scale-free. To the contrary, several important networks do not share this property:

- Networks appearing in material science, describing the bonds between the atoms in crystalline or amorphous materials. In these networks each node has exactly the same degree, determined by chemistry (Figure 4.11).
- The neural network of the C. elegans worm [28].
- The power grid, consisting of generators and switches connected by transmission lines.

For the scale-free property to emerge the nodes need to have the capacity to link to an arbitrary number of other nodes. These links do not need to be concurrent: We do not constantly chat with each of our acquaintances and a protein in the cell does not simultaneously bind to each of its potential interaction partners. The scale-free property is absent in systems that limit the number of links a node can have, effectively restricting the maximum size of the hubs. Such limitations are common in materials (Figure 4.11), explaining why they cannot develop a scale-free topology.

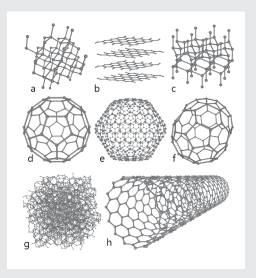


Figure 4.11 The Material Network

A carbon atom can share only four electrons with other atoms, hence no matter how we arrange these atoms relative to each other, in the resulting network a node can never have more than four links. Hence, hubs are forbidden and the scale-free property cannot emerge. The figure shows several carbon allotropes, i.e. materials made of carbon that differ in the structure of the network the carbon atoms arrange themselves in. This different arrangement results in materials with widely different physical and electronic characteristics, like (a) diamond; (b) graphite; (c) lonsdaleite; (d) C60 (buckminsterfullerene); (e) C540 (a fullerene) (f) C70 (another fullerene); (g) amorphous carbon; (h) single-walled carbon nanotube.

ULTRA-SMALL WORLD PROPERTY

The presence of hubs in scale-free networks raises an interesting question: Do hubs affect the small world property? Figure 4.4 suggests that they do: Airlines build hubs precisely to decrease the number of hops between two airports. The calculations support this expectation, finding that distances in a scale-free network are smaller than the distances observed in an equivalent random network.

The dependence of the average distance $\langle d \rangle$ on the system size *N* and the degree exponent γ are captured by the formula [29, 30]

$$\langle d \rangle \sim \begin{cases} \text{const.} & \gamma = 2\\ \ln \ln N & 2 < \gamma < 3\\ \frac{\ln N}{\ln \ln N} & \gamma = 3\\ \ln N & \gamma > 3 \end{cases}$$
(4.22)

Next we discuss the behavior of $\langle d \rangle$ in the four regimes predicted by (4.22), as summarized in Figure 4.12:

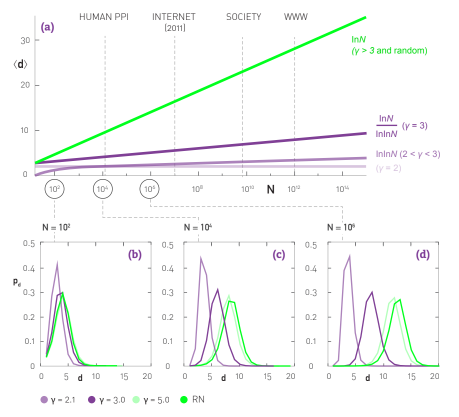
Anomalous Regime (y = 2)

According to (4.18) for $\gamma=2$ the degree of the biggest hub grows linearly with the system size, i.e. $k_{max}\sim N$. This forces the network into a *hub and spoke* configuration in which all nodes are close to each other because they all connect to the same central hub. In this regime the average path length does not depend on N.

Ultra-Small World (2 $< \gamma < 3$)

Equation (4.22) predicts that in this regime the average distance increases as lnln*N*, a significantly slower growth than the ln*N* derived for random networks. We call networks in this regime *ultra-small*, as the hubs radically reduce the path length [29]. They do so by linking to a large number of small-degree nodes, creating short distances between them.

To see the implication of the ultra-small world property consider again the world's social network with $N \approx 7 \times 10^9$. If the society is described by a random network, the *N*-dependent term is $\ln N = 22.66$. In contrast for a scale-free network the *N*-dependent term is $\ln \ln N = 3.12$, indicating that the hubs radically shrink the distance between the nodes.



Critical Point $(\gamma = 3)$

This value is of particular theoretical interest, as the second moment of the degree distribution does not diverge any longer. We therefore call $\gamma = 3$ the *critical point*. At this critical point the $\ln N$ dependence encountered for random networks returns. Yet, the calculations indicate the presence of a double logarithmic correction $\ln \ln N$ [29, 31], which shrinks the distances compared to a random network of similar size.

Small World ($\gamma > 3$)

In this regime $\langle k^2 \rangle$ is finite and the average distance follows the small world result derived for random networks. While hubs continue to be present, for $\gamma > 3$ they are not sufficiently large and numerous to have a significant impact on the distance between the nodes.

Taken together, (4.22) indicates that the more pronounced the hubs are, the more effectively they shrink the distances between nodes. This conclusion is supported by Figure 4.12a, which shows the scaling of the average path length for scale-free networks with different γ . The figure indicates that while for small N the distances in the four regimes are comparable, for large N we observe remarkable differences.

Further support is provided by the path length distribution for scale-

Figure 4.12
Distances in Scale-free Networks

(a) The scaling of the average path length in the four scaling regimes characterizing a scale-free network: constant ($\gamma = 2$), lnlnN (2 < γ < 3), lnN/lnlnN (γ = 3), lnN $(\gamma > 3 \text{ and random networks})$. The dotted lines mark the approximate size of several real networks. Given their modest size, in biological networks, like the human protein-protein interaction network (PPI), the differences in the node-to-node distances are relatively small in the four regimes. The differences in $\langle d \rangle$ is quite significant for networks of the size of the social network or the WWW. For these the small-world formula significantly underestimates the real $\langle d \rangle$.

(b) (c) (d)

Distance distribution for networks of size $N=10^2$, 10^4 , 10^6 , illustrating that while for small networks $(N=10^2)$ the distance distributions are not too sensitive to γ , for large networks $(N=10^6)$ p_d and $\langle d \rangle$ change visibly with γ .

The networks were generated using the static model [32] with $\langle k \rangle$ = 3.

free networks with different γ and N (Figure 4.12b-d). For $N=10^2$ the path length distributions overlap, indicating that at this size differences in γ result in undetectable differences in the path length. For $N=10^6$, however, p_d observed for different γ are well separated. Figure 4.12d also shows that the larger the degree exponent, the larger are the distances between the nodes.

In summary the scale-free property has several effects on network distances:

- Shrinks the average path lengths. Therefore most scale-free networks of practical interest are not only "small", but are "ultra-small". This is a consequence of the hubs, that act as bridges between many small degree nodes.
- Changes the dependence of $\langle d \rangle$ on the system size, as predicted by (4.22). The smaller is γ , the shorter are the distances between the nodes.
- Only for $\gamma > 3$ we recover the $\ln N$ dependence, the signature of the small-world property characterizing random networks (Figure 4.12).

BOX 4.4

WE ARE ALWAYS CLOSE TO THE HUBS

Frigyes Karinthy in his 1929 short story [33] that first described the small world concept cautions that "it's always easier to find someone who knows a famous or popular figure than some runthe-mill, insignificant person". In other words, we are typically closer to hubs than to less connected nodes. This effect is particularly pronounced in scale-free networks (Figure 4.13).

The implications are obvious: There are always short paths linking us to famous individuals like well known scientists or the president of the United States, as they are hubs with an exceptional number of acquaintances. It also means that many of the shortest paths go through these hubs.

In contrast to this expectation, measurements aiming to replicate the six degrees concept in the online world find that individuals involved in chains that reached their target were less likely to send a message to a hub than individuals involved in incomplete chains [34]. The reason may be self-imposed: We perceive hubs as being busy, so we contact them only in real need. We therefore avoid them in online experiments of no perceived value to them.

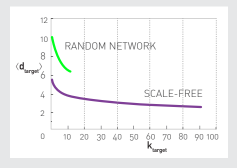


Figure 4.13 Closing on the hubs

The distance $\langle d_{target} \rangle$ of a node with degree $k \approx \langle k \rangle$ to a target node with degree k_{target} in a random and a scale-free network. In scale-free networks we are closer to the hubs than in random networks. The figure also illustrates that in a random network the largest-degree nodes are considerably smaller and hence the path lengths are visibly longer than in a scale-free network. Both networks have $\langle k \rangle = 2$ and N = 1,000 and for the scale-free network we choose $\gamma = 2.5$.

THE ROLE OF THE DEGREE EXPONENT

Many properties of a scale-free network depend on the value of the degree exponent γ . A close inspection of Table 4.1 indicates that:

- γ varies from system to system, prompting us to explore how the properties of a network change with γ .
- For most real systems the degree exponent is above 2, making us wonder: Why don't we see networks with y < 2?

To address these questions next we discuss how the properties of a scale-free network change with γ (BOX 4.5).

Anomalous Regime (γ≤ 2)

For γ < 2 the exponent $1/(\gamma-1)$ in (4.18) is larger than one, hence the number of links connected to the largest hub grows faster than the size of the network. This means that for sufficiently large N the degree of the largest hub must exceed the total number of nodes in the network, hence it will run out of nodes to connect to. Similarly, for γ < 2 the average degree $\langle k \rangle$ diverges in the $N \to \infty$ limit. These odd predictions are only two of the many anomalous features of scale-free networks in this regime. They are signatures of a deeper problem: Large scale-free network with γ < 2, that lack multi-links, cannot exist (BOX 4.6).

Scale-Free Regime $(2 < \gamma < 3)$

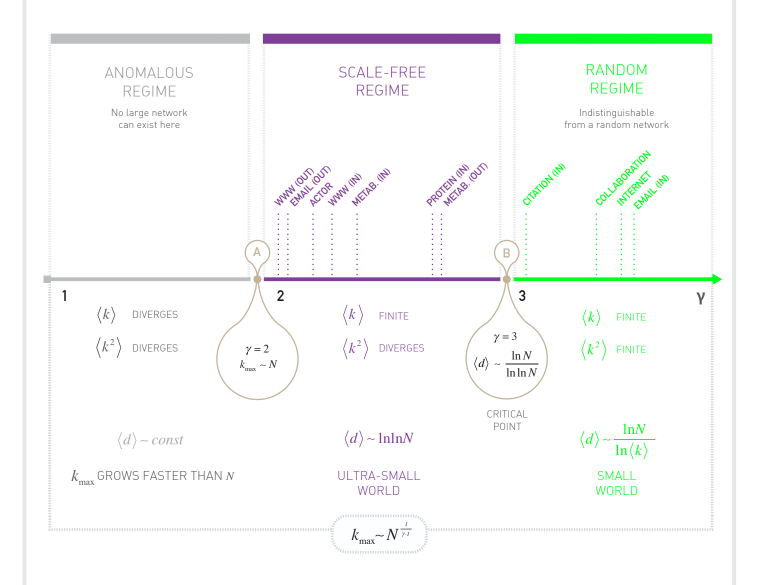
In this regime the first moment of the degree distribution is finite but the second and higher moments diverge as $N \to \infty$. Consequently scale-free networks in this regime are ultra-small (SECTION 4.6). Equation (4.18) predicts that k_{max} grows with the size of the network with exponent 1/ (γ - 1), which is smaller than one. Hence the market share of the largest hub, k_{max} /N, representing the fraction of nodes that connect to it, decreases as k_{max} / $N \sim N^{-(\gamma-2)/(\gamma-1)}$.

As we will see in the coming chapters, many interesting features of scale-free networks, from their robustness to anomalous spreading

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BOX 4.5

THE γ DEPENDENT PROPERTIES OF SCALE-FREE NETWORKS



phenomena, are linked to this regime.

Random Network Regime (y > 3)

According to (4.20) for $\gamma > 3$ both the first and the second moments are finite. For all practical purposes the properties of a scale-free network in this regime are difficult to distinguish from the properties a random network of similar size. For example (4.22) indicates that the average distance between the nodes converges to the small-world formula derived for random networks. The reason is that for large γ the degree distribution p_k decays sufficiently fast to make the hubs small and less numerous.

Note that scale-free networks with large γ are hard to distinguish from a random network. Indeed, to document the presence of a power-law degree distribution we ideally need 2-3 orders of magnitude of scaling, which means that k_{max} should be at least 10^2 - 10^3 times larger than k_{min} . By inverting (4.18) we can estimate the network size necessary to observe the desired scaling regime, finding

$$N = \left(\frac{k_{\text{max}}}{k_{\text{min}}}\right)^{\gamma - 1}.$$
(4.23)

For example, if we wish to document the scale-free nature of a network with $\gamma=5$ and require scaling that spans at least two orders of magnitudes (e.g. $k_{min}\sim 1$ and $k_{max}\simeq 10^2$), according to (4.23) the size of the network must exceed $N>10^8$. There are very few network maps of this size. Therefore, there may be many networks with large degree exponent. Given, however, their limited size, it is difficult to obtain convincing evidence of their scale-free nature.

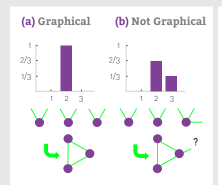
In summary, we find that the behavior of scale-free networks is sensitive to the value of the degree exponent γ . Theoretically the most interesting regime is $2<\gamma<3$, where $\langle k^2\rangle$ diverges, making scale-free networks ultra-small. Interestingly, many networks of practical interest, from the WWW to protein interaction networks, are in this regime.

BOX 4.6

WHY SCALE-FREE NETWORKS WITH γ < 2 DO NOT EXIST

To see why networks with $\gamma < 2$ are problematic, we need to attempt to build one. A degree sequence that can be turned into simple graph (i.e. a graph lacking multi-links or self-loops) is called graphical [35]. Yet, not all degree sequences are graphical: For example, if the number of stubs is odd, then we will always have an unmatched stub (Figure 4.14b).

The graphicality of a degree sequence can be tested with an algorithm proposed by Erdős and Gallai [35, 36, 37, 38, 39]. If we apply the algorithm to scale-free networks we find that the number of graphical degree sequences drops to zero for $\gamma < 2$ (Figure 4.14c). Hence degree distributions with $\gamma < 2$ cannot be turned into simple networks. Indeed, for networks in this regime the largest hub grows faster than N. If we do not allow self-loops and multi-links, then the largest hub will run out of nodes to connect to once its degree exceeds N-1.



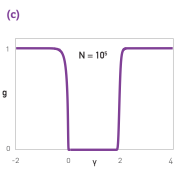


Figure 4.14 Networks With γ < 2 are Not Graphical

- (a-b) Degree distributions and the corresponding degree sequences for two small networks. The difference between them is in the degree of a single node. While we can build a simple network using the degree distribution (a), it is impossible to build one using (b), as one stub always remains unmatched. Hence (a) is graphical, while (b) is not.
- (c) Fraction of networks, g, for a given γ that are graphical. A large number of degree sequences with degree exponent γ and $N=10^5$ were generated, testing the graphicality of each network. The figure indicates that while virtually all networks with $\gamma > 2$ are graphical, it is impossible to find graphical networks in the $0 < \gamma < 2$ range. After [39].

GENERATING NETWORKS WITH ARBITRARY DEGREE DISTRIBUTION

Networks generated by the Erdős-Rényi model have a Poisson degree distribution. The empirical results discussed in this chapter indicate, however, that the degree distribution of real networks significantly deviates from a Poisson form, raising an important question: How do we generate networks with an arbitrary $p_{\rm k}$? In this section we discuss three frequently used algorithms designed for this purpose.

Configuration Model

The configuration model, described in Figure 4.15, helps us build a network with a pre-defined degree sequence. In the network generated by the model each node has a pre-defined degree k_i , but otherwise the network is wired randomly. Consequently the network is often called a *random network with a pre-defined degree sequence*. By repeatedly applying this procedure to the same degree sequence we can generate different networks with the same p_k (Figure 4.15b-d). There are a couple of caveats to consider:

• The probability to have a link between nodes of degree k, and k, is

$$p_{ij} = \frac{k_i k_j}{2L - 1} \tag{4.24}$$

Indeed, a stub starting from node i can connect to 2L-1 other stubs. Of these, k_j are attached to node j. So the probability that a particular stub is connected to a stub of node j is k_j /(2L-1). As node i has k_i stubs, it has k_i attempts to link to j, resulting in (4.24).

• The obtained network contains self-loops and multi-links, as there is nothing in the algorithm to forbid a node connecting to itself, or to generate multiple links between two nodes. We can choose to reject stub pairs that lead to these, but if we do so, we may not be able to complete the network. Rejecting self-loops or multi-links also means that not all possible matchings appear with equal probability. Hence (4.24) will not be valid, making analytical calculations difficult. Yet, the number of self-loops and multi-links remain negligible, as the number of choices to connect to increases with *N*, so typically we do not need to exclude them [42].

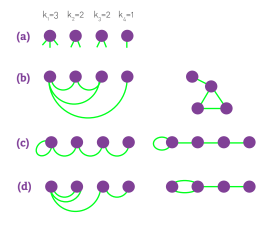


Figure 4.15
The Configuration Model

The configuration model builds a network whose nodes have pre-defined degrees [40, 41]. The algorithm consists of the following steps:

(a) Degree Sequence

Assign a degree to each node, represented as stubs or half-links. The degree sequence is either generated analytically from a preselected p_k distribution (BOX 4.7), or it is extracted from the adjacency matrix of a real network. We must start from an even number of stubs, otherwise we are left with unpaired stubs.

(b, c, d) Network Assembly

Randomly select a stub pair and connect them. Then randomly choose another pair from the remaining 2L - 2 stubs and connect them. This procedure is repeated until all stubs are paired up. Depending on the order in which the stubs were chosen, we obtain different networks. Some networks include cycles (b), others self-loops (c) or multi-links (d). Yet, the expected number of self-loops and multi-links goes to zero in the $N \to \infty$ limit.

• The configuration model is frequently used in calculations, as (4.24) and its inherently random character helps us analytically calculate numerous network measures.

BOX 4.7

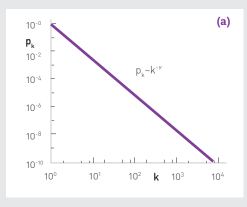
GENERATING A DEGREE SEQUENCE WITH POWER-LAW DISTRIBUTION

The *degree sequence* of an undirected network is a sequence of node degrees. For example, the degree sequence of each of the networks shown in Figure 4.15a is {3, 2, 2, 1}. As Figure 4.15a illustrates, the degree sequence does not uniquely identify a graph, as there are multiple ways we can pair up the stubs.

To generate a degree sequence from a pre-defined degree distribution we start from an analytically pre-defined degree distribution, like $p_k \sim k^{\gamma}$, shown in Figure 4.16a. Our goal is to generate a degree sequence $\{k_1, k_2, ..., k_N\}$ that follow the distribution p_k . We start by calculating the function

$$D(k) = \sum_{k \ge k} p_{k'} \tag{4.25}$$

shown in Figure 4.16b. D(k) is between 0 and 1, and the step size at any k equals p_k . To generate a sequence of N degrees following p_k , we generate N random numbers r_i , i=1,...,N, chosen uniformly from the (0,1) interval. For each r_i we use the plot in (b) to assign a degree k_i . The obtained $k_i = D^{-1}(r_i)$ set of numbers follows the desired p_k distribution. Note that the degree sequence assigned to a p_k is not unique - we can generate multiple sets of $\{k_1,...,k_N\}$ sequences compatible with the same p_k .



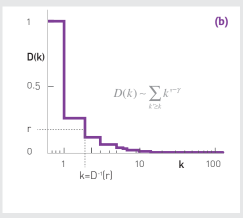


Figure 4.16 Generating a Degree Sequence

- (a) The power law degree distribution of the degree sequence we wish to generate.
- **(b)** The function (4.25), that allows us to assign degrees k to uniformly distributed random numbers r.

Degree-Preserving Randomization

As we explore the properties of a real network, we often need to ask if a certain network property is predicted by its degree distribution alone, or if it represents some additional property not contained in p_k . To answer this question we need to generate networks that are wired randomly, but whose p_k is identical to the original network. This can be achieved through degree-preserving randomization [43] described in Figure 4.17b. The idea behind the algorithm is simple: We randomly select two links and swap them, if the swap does not lead to multi-links. Hence the degree of each of the four involved nodes in the swap remains unchanged. Consequently, hubs stay hubs and small-degree nodes retain their small degree, but the wiring diagram of the generated network is randomized. Note that degree-preserving randomization is different from full randomization, where we swap links without preserving the node degrees (Figure 4.17a). Full randomization turns any network into an Erdős-Rényi network with a Poisson degree distribution that is independent of the original p_k .

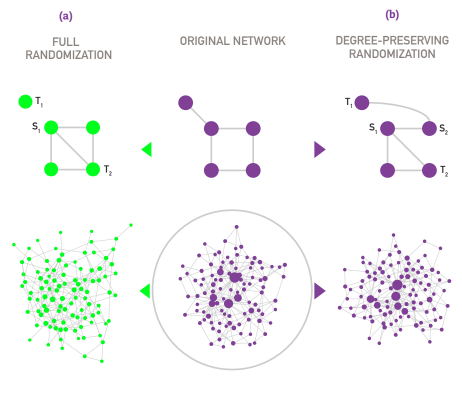


Figure 4.17

Degree Preserving Randomization

Two algorithms can generate a randomized version of a given network [43], with different outcomes.

(a) Full Randomization

This algorithm generates a random (Erdős-Rényi) network with the same N and L as the original network. We select randomly a source node (S_1) and two target nodes, where the first target (T_1) is linked directly to the source node and the second target (T_2) is not. We rewire the S_1 - T_1 link, turning it into an S_1 - T_2 link. As a result the degree of the target nodes T_1 and T_2 changes. We perform this procedure once for each link in the network.

(b) Degree-Preserving Randomization

This algorithm generates a network in which each node has exactly the same degree as in the original network, but the network's wiring diagram has been randomized. We select two source (S_1, S_2) and two target nodes (T_1, T_2) , such that initially there is a link between S_1 and T_1 , and a link between S_2 and T_2 . We then swap the two links, creating an S_1 - T_2 and an S_2 - T_1 link. The swap leaves the degree of each node unchanged.We repeat this procedure until we rewire each link at least once.

Bottom Panels: Starting from a scale-free network (middle), full randomization eliminates the hubs and turns the network into a random network (left). In contrast, degree-preserving randomization leaves the hubs in place and the network remains scale-free (right).

Hidden Parameter Model

The configuration model generates self-loops and multi-links, features that are absent in many real networks. We can use the *hidden parameter model* (Figure 4.18) to generate networks with a pre-defined p_k but without multi-links and self-loops [44, 45, 46].

We start from N isolated nodes and assign each node i a hidden parameter η_i , chosen from a distribution $\rho(\eta)$. The nature of the generated network depends on the selection of the $\{\eta_i\}$ hidden parameter sequence. There are two ways to generate the appropriate hidden parameters:

• η_i can be a sequence of N random numbers chosen from a pre-defined $\rho(\eta)$ distribution. The degree distribution of the obtained network is

$$p_{k} = \int \frac{e^{-\eta} \eta^{k}}{k!} \rho(\eta) d\eta. \tag{4.26}$$

• η_i can come from a deterministic sequence $\{\eta_1, \eta_2, ..., \eta_N\}$. The degree distribution of the obtained network is

$$p_{k} = \frac{1}{N} \sum_{j} \frac{e^{-\eta_{j}} \eta_{j}^{k}}{k!}.$$
 (4.27)

The hidden parameter model offers a particularly simple method to generate a scale-free network. Indeed, using

$$\eta_j = \frac{c}{i^{\alpha}}, i = 1, \dots, N \tag{4.28}$$

as the sequence of hidden parameters, according to (4.27) the obtained network will have the degree distribution

$$p_{k} \sim k^{-(1+\frac{1}{\alpha})}$$
 (4.29)

for large k. Hence by choosing the appropriate α we can tune $\gamma=1+1/\alpha$. We can also use $\langle \eta \rangle$ to tune $\langle k \rangle$ as (4.26) and (4.27) imply that $\langle k \rangle = \langle \eta \rangle$.

In summary, the configuration model, degree-preserving randomization and the hidden parameter model can generate networks with a pre-defined degree distribution and help us analytically calculate key network characteristics. We will turn to these algorithms each time we explore whether a certain network property is a consequence of the network's degree distribution, or if it represents some emergent property (BOX 4.8). As we use these algorithms, we must be aware of their limitations:

- The algorithms do not tell us why a network has a certain degree distribution. Understanding the origin of the observed p_k will be the subject of CHAPTERS 6 and 7.
- Several important network characteristics, from clustering (CHAPTER 9) to degree correlations (CHAPTER 7), are lost during randomization.

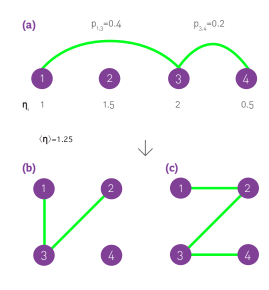


Figure 4.18 Hidden Parameter Model

(a) We start with N isolated nodes and assign to each node a *hidden parameter* η_i , which is either selected from a $\rho(\eta)$ distribution or it is provided by a sequence $\{\eta_i\}$. We connect each node pair with probability

$$p(\eta_i,\eta_j) = \frac{\eta_i \eta_j}{\langle \eta \rangle N}.$$

The figure shows the probability to connect nodes (1,3) and (3,4).

(b, c) After connecting the nodes, we obtain the networks shown in (b) or (c), representing two independent realizations generated by the same hidden parameter sequence (a).

The expected number of links in the network generated by the model is

$$L = \frac{1}{2} \sum_{i,j'}^{N} \frac{\eta_i \eta_j}{\langle \eta \rangle N} = \frac{1}{2} \langle \eta \rangle N.$$

Similar to the random network model, L will vary from network to network, following an exponentially bounded distribution. If we wish to control the average degree $\langle k \rangle$ we can add L links to the network one by one. The end points i and j of each link are then chosen randomly with a probability proportional to η_i and η_j . In this case we connect i and j only if they were not connected previously.

BOX 4.8

TESTING THE SMALL-WORD PROPERTY

In the literature the distances observed in a real network are often compared to the small-world formula (3.19). Yet, (3.19) was derived for random networks, while real networks do not have a Poisson degree distribution. If the network is scale-free, then (4.22) offers the appropriate formula. Yet, (4.22) provides only the scaling of the distance with N, and not its absolute value. Instead of fitting the average distance, we often ask: Are the distances observed in a real network comparable with the distances observed in a randomized network with the same degree distribution? Degree preserving randomization helps answer this question. We illustrate the procedure on the protein interaction network.

(i) Original Network

We start by measuring the distance distribution p_d of the original network, obtaining $\langle d \rangle$ = 5.61 (Figure 4.19).

(ii) Full Randomization

We generate a random network with the same N and L as the original network. The obtained p_d visibly shifts to the right, providing $\langle d \rangle = 7.13$, much larger than the original $\langle d \rangle = 5.61$. It is tempting to conclude that the protein interaction network is affected by some unknown organizing principle that keeps the distances shorter. This would be a flawed conclusion, however, as the bulk of the difference is due to the fact that full randomization changed the degree distribution.

(iii) Degree-Preserving Randomization

As the original network is scale-free, the proper random reference should maintain the original degree distribution. Hence we determine $\boldsymbol{p}_{\scriptscriptstyle d}$ after degree-preserving randomization, finding that it is comparable to the original $\boldsymbol{p}_{\scriptscriptstyle d}$.

In summary, a random network overestimates the distances between the nodes, as it is missing the hubs. The network obtained by degree preserving randomization retains the hubs, so the distances of the randomized network are comparable to the original network. This example illustrates the importance of choosing the proper randomization procedure when exploring networks.

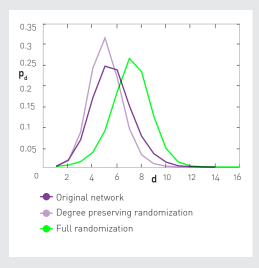


Figure 4.19 Randomizing Real Networks

The distance distribution p_d between each node pair in the protein-protein interaction network (Table 4.1). The green line provides the path-length distribution obtained under *full randomization*, which turns the network into an Erdős-Rényi network, while keeping N and L unchanged (Figure 4.17).

The light purple curve correspond to p_d of the network obtained after *degree-preserving randomization*, which keeps the degree of each node unchanged.

We have: $\langle d \rangle$ =5.61±1.64 (original), $\langle d \rangle$ =7.13 ± 1.62 (full randomization), $\langle d \rangle$ =5.08 ± 1.34 (degree-preserving randomization).

Hence, the networks generated by these algorithms are a bit like a photograph of a painting: at first look they appear to be the same as the original. Upon closer inspection we realize, however, that many details, from the texture of the canvas to the brush strokes, are lost.

The three algorithms discussed above raise the following question: How do we decide which one to use? Our choice depends on whether we start from a degree sequence $\{k_i\}$ or a degree distribution p_k and whether we can tolerate self-loops and multi-links between two nodes. The decision tree involved in this choice is provided in Figure 4.20.

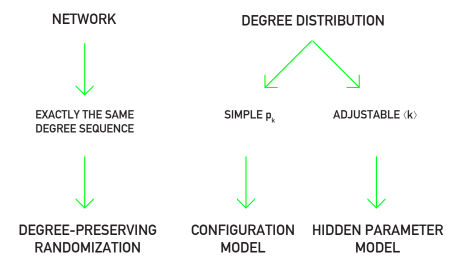


Figure 4.20
Choosing a Generative Algorithm

The choice of the appropriate generative algorithm depends on several factors. If we start from a real network or a known degree sequence, we can use *degree-preserving randomization*, which guarantees that the obtained networks are simple and have the degree sequence of the original network. The model allows us to forbid multi-links or self-loops, while maintaining the degree sequence of the original network.

If we wish to generate a network with given pre-defined degree distribution $p_{\mathbf{k}}$, we have two options. If $p_{\mathbf{k}}$ is known, the configuration model offers a convenient algorithm for network generation. For example, the model allows us generate a networks with a pure power law degree distribution $p_{\mathbf{k}} = \mathsf{C} k^{-\gamma}$ for $k \geq k_{\min}$.

However, tuning the average degree $\langle k \rangle$ of a scale-free network within the configuration model is a tedious task, because the only available free parameter is k_{\min} . Therefore, if we wish to alter $\langle k \rangle$, it is more convenient to use the hidden parameter model with parameter sequence (4.28). This way the tail of the degree distribution follows $\sim k^{-\gamma}$ and by changing the number of links L we can to control $\langle k \rangle$.

SUMMARY

The scale-free property has played an important role in the development of network science for two main reasons:

- Many networks of scientific and practical interest, from the WWW to the subcellular networks, are scale-free. This universality made the scale-free property an unavoidable issue in many disciplines.
- Once the hubs are present, they fundamentally change the system's behavior. The ultra-small property offers a first hint of their impact on a network's properties; we will encounter many more examples in the coming chapters.

As we continue to explore the consequences of the scale-free property, we must keep in mind that the power-law form (4.1) is rarely seen in this pure form in real systems. The reason is that a host of processes affect the topology of each network, which also influence the shape of the degree distribution. We will discuss these processes in the coming chapters. The diversity of these processes and the complexity of the resulting \boldsymbol{p}_k confuses those who approach these networks through the narrow perspective of the quality of fit to a pure power law. Instead the scale-free property tells us that we must distinguish two rather different classes of networks:

Exponentially Bounded Networks

We call a network *exponentially bounded* if its degree distribution decrease exponentially or faster for high k. As a consequence $< k^2 >$ is smaller than < k >, implying that we lack significant degree variations. Examples of p_k in this class include the Poisson, Gaussian, or the simple exponential distribution (Table 4.2). Erdős-Rényi and Watts-Strogatz networks are the best known models network belonging to this class. Exponentially bounded networks lack outliers, consequently most nodes have comparable degrees. Real networks in this class include highway networks and the power grid.

Fat Tailed Networks

We call a network *fat tailed* if its degree distribution has a power law tail in the high-k region. As a consequence $< k^2 >$ is much larger than < k >, resulting in considerable degree variations. Scale-free networks with a power-law degree distribution (4.1) offer the best known example of networks belonging to this class. Outliers, or exceptionally high-degree

nodes, are not only allowed but are expected in these networks. Networks in this class include the WWW, the Internet, protein interaction networks, and most social and online networks.

While it would be desirable to statistically validate the precise form of the degree distribution, often it is sufficient to decide if a given network has an exponentially bounded or a fat tailed degree distribution (see ADVANCED TOPICS 4.A). If the degree distribution is exponentially bounded, the random network model offers a reasonable starting point to understand its topology. If the degree distribution is fat tailed, a scale-free network offers a better approximation. We will also see in the coming chapters that the key signature of the fat tailed behavior is the magniture of $\langle k^2 \rangle$: If $\langle k^2 \rangle$ is large, systems behave like scale-free networks; if $\langle k^2 \rangle$ is small, being comparable to $\langle k \rangle (\langle t \rangle + 1)$, systems are well approximated by random networks.

In summary, to understand the properties of real networks, it is often sufficient to remember that in scale-free networks a few highly connected hubs coexist with a large number of small nodes. The presence of these hubs plays an important role in the system's behavior. In this chapter we explored the basic characteristics of scale-free networks. We are left, therefore, with an important question: Why are so many real networks scale-free? The next chapter provides the answer.

BOX 4.9

AT A GLANCE: SCALE-FREE NETWORKS

DEGREE DISTRIBUTION

Discrete form:

$$p_k = \frac{k^{-\gamma}}{\zeta(\gamma)}$$

Continuous form:

$$p(k) = (\gamma - 1)k_{min}^{\gamma - 1} k^{-\gamma}$$

SIZE OF THE LARGEST HUB

$$k_{\max} = k_{\min} N^{\frac{1}{y-1}}.$$

MOMENTS OF p_k for $N \to \infty$ 2 < $\gamma \le 3$: $\langle k \rangle$ finite, $\langle k^2 \rangle$ diverges.

 $\gamma > 3$: $\langle k \rangle$ and $\langle k^2 \rangle$ finite.

DISTANCES

$$\langle d \rangle \sim \begin{cases} \text{const.} & \gamma = 2\\ \ln \ln N & 2 < \gamma < 3\\ \frac{\ln N}{\ln \ln N} & \gamma = 3\\ \ln N & \gamma > 3 \end{cases}$$

HOMEWORK

4.1. Hubs

Calculate the expected maximum degree $k_{\rm max}$ for the undirected networks listed in Table 4.1.

4.2. Friendship Paradox

The degree distribution p_k expresses the probability that a randomly selected node has k neighbors. However, if we randomly select a link, the probability that a node at one of its ends has degree k is $q_k = Akp_k$, where A is a normalization factor.

- (a) Find the normalization factor A, assuming that the network has a power law degree distribution with $2 < \gamma < 3$, with minimum degree k_{\min} and maximum degree k_{\max} .
- (b) In the configuration model q_k is also the probability that a randomly chosen node has a neighbor with degree k. What is the average degree of the neighbors of a randomly chosen node?
- (c) Calculate the average degree of the neighbors of a randomly chosen node in a network with $N=10^4$, $\gamma=2.3$, $k_{\rm min}=1$ and $k_{\rm max}=1$, 000. Compare the result with the average degree of the network, $\langle k \rangle$.
- (d) How can you explain the "paradox" of (c), that is a node's friends have more friends than the node itself?

4.3. Generating Scale-Free Networks

Write a computer code to generate networks of size N with a power-law degree distribution with degree exponent γ . Refer to SECTION 4.9 for the procedure. Generate three networks with $\gamma=2.2$ and with $N=10^3$, $N=10^4$ and $N=10^5$ nodes, respectively. What is the percentage of multi-link and self-loops in each network? Generate more networks to plot this percentage in function of N. Do the same for networks with $\gamma=3$.

4.4. Mastering Distributions

Use a software which includes a statistics package, like Matlab, Math-

ematica or Numpy in Python, to generate three synthetic datasets, each containing 10,000 integers that follow a power-law distribution with γ = 2.2, γ = 2.5 and γ = 3. Use $k_{\rm min}$ = 1. Apply the techniques described in AD-VANCED TOPICS 4.C to fit the three distributions.

ADVANCED TOPICS 4.A POWER LAWS

Power laws have a convoluted history in natural and social sciences, being interchangeably (and occasionally incorrectly) called *fat-tailed*, *heavytailed*, *long-tailed*, *Pareto*, or *Bradford distributions*. They also have a series of close relatives, like *log-normal*, *Weibull*, or *Lévy distributions*. In this section we discuss some of the most frequently encountered distributions in network science and their relationship to power laws.

Exponentially Bounded Distributions

Many quantities in nature, from the height of humans to the probability of being in a car accident, follow bounded distributions. A common property of these is that p_x decays either exponentially (e^{-x}) , or faster than exponentially (e^{-x^2/σ^2}) for high x. Consequently the largest expected x is bounded by some upper value x_{max} that is not too different from $\langle x \rangle$. Indeed, the expected largest x obtained after we draw N numbers from a bounded p_x grows as $x_{max} \sim \log N$ or slower. This means that outliers, representing unusually high x-values, are rare. They are so rare that they are effectively forbidden, meaning that they do not occur with any meaningful probability. Instead, most events drawn from a bounded distribution are in the vicinity of $\langle x \rangle$.

The high-x regime is called the *tail of a distribution*. Given the absence of numerous events in the tail, these distributions are also called *thin tailed*.

Analytically the simplest bounded distribution is the exponential distribution e^{-Ax}. Within network science the most frequently encountered bounded distribution is the Poisson distribution (or its parent, the binomial distribution), which describes the degree distribution of a random network. Outside network science the most frequently encountered member of this class is the normal (Gaussian) distribution (Table 4.2).

Fat Tailed Distributions

The terms fat tailed, heavy tailed, or long tailed refer to p_x whose decay

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at large x is slower than exponential. In these distributions we often encounter events characterized by very large x values, usually called *outliers* or *rare events*. The power-law distribution (4.1) represents the best known example of a fat tailed distribution. An instantly recognizable feature of an fat tailed distribution is that the magnitude of the events x drawn from it can span several orders of magnitude. Indeed, in these distributions the size of the largest event after x0 trials scales as x1 where x2 is determined by the exponent x3 characterizing the tail of the x4 distribution. As x5 grows fast, rare events or outliers occur with a noticeable frequency, often dominating the properties of the system.

The relevance of fat tailed distributions to networks is provided by several factors:

- Many quantities occurring in network science, like degrees, link weights and betweenness centrality, follow a power-law distribution in both real and model networks.
- The power-law form is analytically predicted by appropriate network models (CHAPTER 5).

Crossover Distribution (Log-Normal, Stretched Exponential)

When an empirically observed distribution appears to be between a power law and exponential, *crossover distributions* are often used to fit the data. These distributions may be exponentially bounded (power law with exponential cutoff), or not bounded but decay faster than a power law (log-normal or stretched exponential). Next we discuss the properties of several frequently encountered crossover distributions.

Power law with exponential cut-off is often used to fit the degree distribution of real networks. Its density function has the form:

$$p(x) = Cx^{-\gamma} e^{-\lambda x}, \tag{4.30}$$

$$C = \frac{\lambda^{1-\gamma}}{\Gamma(1-\gamma, \lambda x_{\min})}$$
 (4.31)

where x > 0 and $\gamma > 0$ and $\Gamma(s,y)$ denotes the upper incomplete gamma function. The analytical form (4.30) directly captures its crossover nature: it combines a power-law term, a key component of fat tailed distributions, with an exponential term, responsible for its exponentially bounded tail. To highlight its crossover characteristics we take the logarithm of (4.30),

$$\ln p(x) = \ln C - \gamma \ln x - \lambda x. \tag{4.32}$$

For $x \ll 1/\lambda$ the second term on the r.h.s dominates, suggesting that the distribution follows a power law with exponent γ . Once $x \gg 1/\lambda$, the λx term overcomes the ln x term, resulting in an exponential cutoff for high x.

Stretched exponential (Weibull distribution) is formally similar to (4.30) except that there is a fractional power law in the exponential. Its name comes from the fact that its cumulative distribution function is one minus a stretched exponential function $P(x) = e^{-(\lambda x)^{\mu}}$ (4.32) which leads to density function

$$P'(x) = Cx^{\beta - 1} e^{-(\lambda x)^{\beta}}$$
 (4.33)

$$C = \beta \lambda^{\beta}. \tag{4.34}$$

In most applications x varies between 0 and $+\infty$. In (4.32) β is the *stretching exponent*, determining the properties of p(x):

- For $\beta = 1$ we recover a simple exponential function.
- If β is between 0 and 1, the graph of log p(x) versus x is "stretched", meaning that it spans several orders of magnitude in x. This is the regime where a stretched exponential is difficult to distinguish from a pure power law. The closer β is to 0, the more similar is p(x) to the power law x^{-1} .
- If $\beta > 1$ we have a "compressed" exponential function, meaning that x varies in a very narrow range.
- For β = 2 (4.33) reduces to the Rayleigh distribution.

As we will see in CHAPTERS 5 and 6, several network models predict a streched exponential degree distribution.

A log-normal distribution (Galton or Gibrat distribution) emerges if $\ln x$ follows a normal distribution. Typically a variable follows a log-normal distribution if it is the product of many independent positive random numbers. We encounter log-normal distributions in finance, representing the compound return from a sequence of trades.

The probability density function of a log-normal distribution is

$$p(x) = \frac{1}{\sqrt{2\pi}\sigma x} \exp\left[-\frac{(\ln x - \mu)^2}{2\sigma^2}\right]$$
 (4.35)

Hence a log-normal is like a normal distribution except that its variable in the exponential term is not x, but ln x.

To understand why a log-normal is occasionally used to fit a power law distribution, we note that

$$\sigma^{2} = \langle (\ln x)^{2} \rangle - \langle \ln x \rangle^{2}$$
 (4.36)

captures the typical variation of the order of magnitude of x. Therefore now $\ln x$ follows a normal distribution, which means that x can vary rather widely. Depending on the value of σ the log-normal distribution

may resemble a power law for several orders of magnitude. This is also illustrated in Table 4.2, that shows that $\langle x^2 \rangle$ grows exponentially with σ , hence it can be very large.

In summary, in most areas where we encounter fat-tailed distributions, there is an ongoing debate asking which distribution offers the best fit to the data. Frequently encountered candidates include a power law, a stretched exponential, or a log-normal function. In many systems empirical data is not sufficient to distinguish these distributions. Hence as long as there is empirical data to be fitted, the debate surrounding the best fit will never die out.

The debate is resolved by accurate mechanistic models, which analytically predict the expected degree distribution. We will see in the coming chapters that in the context of networks the models predict Poisson, simple exponential, stretched exponential, and power law distributions. The remaining distributions in Table 4.2 are occasionally used to fit the degrees of some networks, despite the fact that we lack theoretical basis for their relevance for networks.

NAME	$p_x/p(x)$	$\langle x \rangle$	$\langle x^2 angle$
Poisson (discrete)	$e^{-\mu}\mu^x/x!$	μ	$\mu(1+\mu)$
Exponential (discrete)	$(1 - e^{-\lambda})e^{-\lambda x}$	$1/(e^{\lambda}-1)$	$(e^{\lambda}+1)/(e^{\lambda}-1)^2$
Exponential (continuous)	$\lambda e^{-\lambda x}$	$1/\lambda$	$2/\lambda^2$
Power law (discrete)	$x^{-\alpha}/\zeta(\alpha)$	$\begin{cases} \zeta(\alpha - 2)/\zeta(\alpha), & \text{if } \alpha > 2\\ \infty, & \text{if } \alpha \le 1 \end{cases}$	$\begin{cases} \zeta(\alpha - 1)/\zeta(\alpha), & \text{if } \alpha > 1\\ \infty, & \text{if } \alpha \le 2 \end{cases}$
Power law (continuous)	$\alpha x^{-\alpha}$	$\begin{cases} \alpha/(\alpha-1), & \text{if } \alpha > 2\\ \infty, & \text{if } \alpha \le 1 \end{cases}$	$\begin{cases} \alpha/(\alpha-2), & \text{if } \alpha > 1\\ \infty, & \text{if } \alpha \le 2 \end{cases}$
Power law with cutoff (continuous)	$\frac{\lambda^{1-\alpha}}{\Gamma(1-\alpha)}x^{-\alpha}e^{-\lambda x}$	$\lambda^{-1} \frac{\Gamma(2-lpha)}{\Gamma(1-lpha)}$	$\lambda^{-2} \frac{\Gamma(3-\alpha)}{\Gamma(1-\alpha)}$
Stretched exponential (continuous)	$\beta \lambda^{\beta} x^{\beta - 1} e^{-(\lambda x)^{\beta}}$	$\lambda^{-1}\Gamma(1+\beta^{-1})$	$\lambda^{-2}\Gamma(1+2\beta^{-1})$
Log-normal (continuous)	$\frac{1}{x\sqrt{2\pi\sigma^2}}e^{-(\ln x - \mu)^2/(2\sigma^2)}$	$e^{\mu+\sigma^2/2}$	$e^{2(\mu+\sigma^2)}$
N ormal (continuous)	$\frac{1}{\sqrt{2\pi\sigma^2}}e^{-(x-\mu)^2/(2\sigma^2)}$	μ	$\mu^2 + \sigma^2$

Table 4.2

Distributions in Network Science

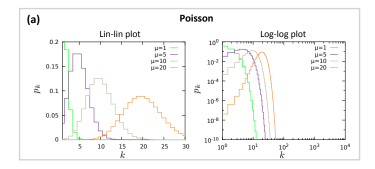
The table lists frequently encountered distributions in network science. For each distribution we show the density function p_x , the appropriate normalization constant C such that

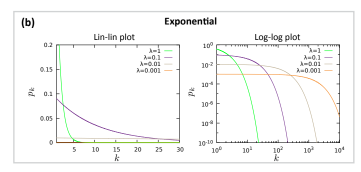
$$\int_{x=x_{\min}}^{\infty} Cf(x) \, dx = 1$$

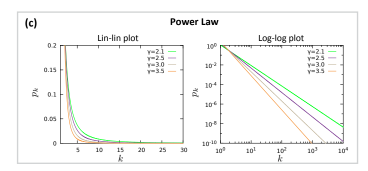
for the continuous case or

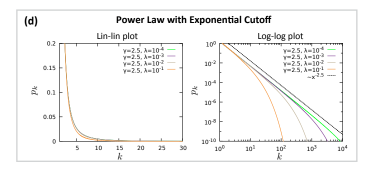
$$\sum_{x=x_{\min}}^{\infty} Cf(x) = 1$$

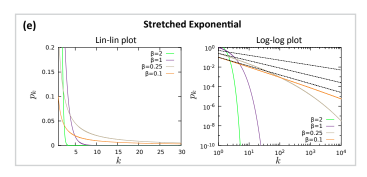
for the discrete case. Given that $\langle x \rangle$ and $\langle x^2 \rangle$ play an important role in network theory, we show the analytical form of these two quantities for each distribution. As some of these distributions diverge at x=0, for most of them $\langle x \rangle$ and $\langle x^2 \rangle$ are calculated assuming that there is a small cutoff x_{\min} in the system. In networks x_{\min} often corresponds to the smallest degree, k_{\min} , or the smallest degree for which the appropriate distribution offers a good fit.

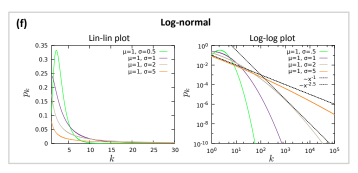












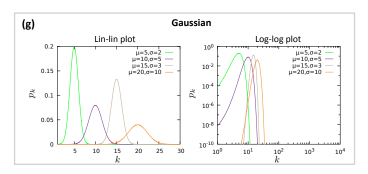


Figure 4.21
Distributions Visualized

Linear and the log-log plots for the most frequently encountered distributions in network science. For definitions see Table 4.2.

THE SCALE-FREE PROPERTY 43 4.A POWER LAWS

ADVANCED TOPICS 4.B PLOTTING POWER-LAWS

Plotting the degree distribution is an integral part of analyzing the properties of a network. The process starts with obtaining N_k , the number of nodes with degree k. This can be provided by direct measurement or by a model. From N_k we calculate $p_k = N_k / N$. The question is, how to plot p_k to best extract its properties.

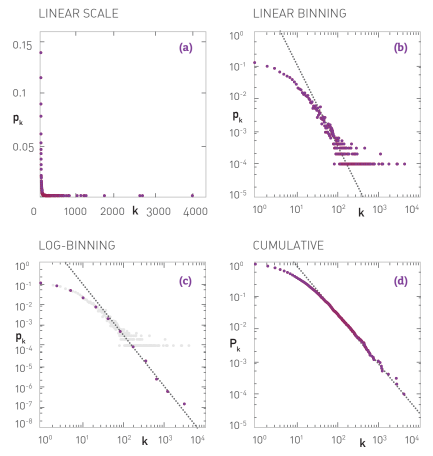
Use a Log-Log Plot

In a scale-free network numerous nodes with one or two links coexist with a few hubs, representing nodes with thousands or even millions of links. Using a linear k-axis compresses the numerous small degree nodes in the small-k region, rendering them invisible. Similarly, as there can be orders of magnitude differences in p_k for k=1 and for large k, if we plot p_k on a linear vertical axis, its value for large k will appear to be zero (Figure 4.22a). The use of a log-log plot avoids these problems. We can either use logarithmic axes, with powers of 10 (used throughout this book, Figure 4.22b) or we can plot $\log p_k$ in function of $\log k$ (equally correct, but slightly harder to read). Note that points with $p_k=0$ or k=0 are not shown on a $\log \log p$ at $\log 0=-\infty$.

Avoid Linear Binning

The most flawed method (yet frequently seen in the literature) is to simply plot $p_k = N_k/N$ on a log-log plot (Figure 4.22b). This is called *linear binning*, as each bin has the same size $\Delta k = 1$. For a scale-free network linear binning results in an instantly recognizable plateau at large k, consisting of numerous data points that form a horizontal line (Figure 4.22b). This plateau has a simple explanation: Typically we have only one copy of each high degree node, hence in the high-k region we either have $N_k = 0$ (no node with degree k) or $N_k = 1$ (a single node with degree k). Consequently linear binning will either provide $p_k = 0$, not shown on a log-log plot, or $p_k = 1/N$, which applies to all hubs, generating a plateau at $p_k = 1/N$.

This plateau affects our ability to estimate the degree exponent γ . For example, if we attempt to fit a power law to the data shown in Figure



4.22b using linear binning, the obtained γ is quite different from the real value γ =2.5. The reason is that under linear binning we have a large number of nodes in small k bins, allowing us to confidently fit p_k in this regime. In the large-k bins we have too few nodes for a proper statistical estimate of p_k . Instead the emerging plateau biases our fit. Yet, it is precisely this high-k regime that plays a key role in determining γ . Increasing the bin size will not solve this problem. It is therefore recommended to avoid linear binning for fat tailed distributions.

Use Logarithmic Binning

Logarithmic binning corrects the non-uniform sampling of linear binning. For log-binning we let the bin sizes increase with the degree, making sure that each bin has a comparable number of nodes. For example, we can choose the bin sizes to be multiples of 2, so that the first bin has size b_0 =1, containing all nodes with k=1; the second has size b_1 =2, containing nodes with degrees k=2, 3; the third bin has size b_2 =4 containing nodes with degrees k=4, 5, 6, 7. By induction the nth bin has size 2^{n-1} and contains all nodes with degrees k=2ⁿ⁻¹, 2ⁿ⁻¹+1, ..., 2ⁿ⁻¹-1. Note that the bin size can increase with arbitrary increments, b_n = c^n , where c > 1. The degree distribution is given by $p_{\langle k_n \rangle} = N_n/b_n$, where N_n is the number of nodes found in the bin n of size b_n and $\langle k_n \rangle$ is the average degree of the nodes in bin b_n .

The logarithmically binned p_k is shown in Figure 4.22c. Note that now the scaling extends into the high-k plateau, invisible under linear binning. Therefore logarithmic binning extracts useful information from the

Figure 4.22

Plotting a Degree Distributions

A degree distribution of the form $p_k \sim (k+k_0)^{-\gamma}$, with k_0 =10 and γ =2.5, plotted using the four procedures described in the text:

(a) Linear Scale, Linear Binning.

It is impossible to see the distribution on a lin-lin scale. This is the reason why we always use log-log plot for scale-free networks.

(b) Log-Log Scale, Linear Binning.

Now the tail of the distribution is visible but there is a plateau in the high-k regime, a consequence of linear binning.

(c) Log-Log Scale, Log-Binning.

With log-binning the plateau dissappears and the scaling extends into the high-*k* regime. For reference we show as light grey the data of (b) with linear binning.

(d) Log-Log Scale, Cumulative.

The cumulative degree distribution shown on a log-log plot.

rare high degree nodes as well (BOX 4.10).

Use Cumulative Distribution

Another way to extract information from the tail of \boldsymbol{p}_k is to plot the complementary cumulative distribution

$$P_k = \sum_{q=k+1}^{\infty} p_q \,, \tag{4.37}$$

which again enhances the statistical significance the high-degree region. If $p_{\rm k}$ follows the power law (4.1), then the cumulative distribution scales as

$$P_{k} \sim k^{-\gamma+1}.\tag{4.38}$$

The cumulative distribution again eliminates the plateau observed for linear binning and leads to an extended scaling region (Figure 4.22d), allowing for a more accurate estimate of the degree exponent.

In summary, plotting the degree distribution to extract its features requires special attention. Mastering the appropriate tools can help us better explore the properties of real networks (80X 4.10).

BOX 4.10

DEGREE DISTRIBUTION OF REAL NETWORKS

In real systems we rarely observe a degree distribution that follows a pure power law. Instead, for most real systems p_k has the shape shown in Figure 4.23a, with some recurring features:

- Low-degree saturation is a common deviation from the power-law behavior. Its signature is a flattened p_k for $k < k_{\rm sat}$. This indicates that we have fewer small degree nodes than expected for a pure power law. The origin of the saturation will be explained in CHAPTER 6.
- High-degree cutoff appears as a rapid drop in p_k for $k > k_{\rm cut}$, indicating that we have fewer high-degree nodes than expected in a pure power law. This limits the size of the largest hub, making it smaller than predicted by (4.18). High-degree cutoffs emerge if there are inherent limitations in the number of links a node can have. For example, in social networks individuals have difficulty maintaining meaningful relationships with an exceptionally large number of acquaintances.

Given the widespread presence of such cutoffs the degree distribution is occasionally fitted to

$$p_x = a(k + k_{\text{sat}})^{-\gamma} \exp\left(-\frac{k}{k_{\text{cut}}}\right).$$
 (4.39)

where $k_{\rm sat}$ accounts for degree saturation, and the exponential term accounts for high-k cutoff. To extract the full extent of the scaling we plot

$$\tilde{p}_x = p_x \exp\left(\frac{k}{k_{\text{cut}}}\right) \tag{4.40}$$

in function of $\tilde{k} = k + k_{\text{sat}}$. According to (4.40) $\tilde{p} \sim \tilde{k}^{-\gamma}$, correcting for the two cutoffs, as seen in Figure 4.23b.

It is occasionally claimed that the presence of low-degree or high-degree cutoffs implies that the network is not scale-free. This is a misunderstanding of the scale-free property: Virtually all properties of scale-free networks are insensitive to the low-degree saturation. Only the high-degree cutoff affects the system's properties by limiting the divergence of the second moment, $\langle k^2 \rangle$. The presence of such cutoffs indicates the presence of additional phenomena that need to be understood.

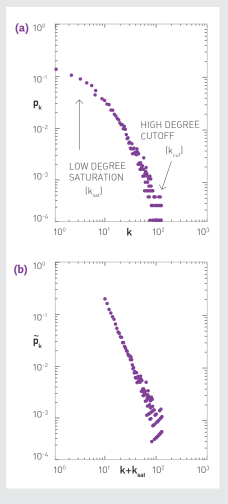


Figure 4.23
Rescaling the Degree Distribution

- (a) In real networks the degree distribution frequently deviates from a pure power law by showing a low degree saturation and high degree cutoff.
- **(b)** By plotting the rescaled P_k in function of $(k + k_{sat})$, as suggested by **(4.40)**, the degree distribution follows a power law for all degrees.

ADVANCED TOPICS 4.C ESTIMATING THE DEGREE EXPONENT

As the properties of scale-free networks depend on the degree exponent (SECTION 4.7), we need to determine the value of γ . We face several difficulties, however, when we try to fit a power law to real data. The most important is the fact that the scaling is rarely valid for the full range of the degree distribution. Rather we observe small- and high- degree cutoffs (BOX 4.10), denoted in this section with K_{\min} and K_{\max} , within which we have a clear scaling region. Note that K_{\min} and K_{\max} are different from k_{\min} and k_{\max} , the latter corresponding to the smallest and largest degrees in a network. They can be the same as k_{sat} and k_{cut} discussed in BOX 4.10. Here we focus on estimating the small degree cutoff K_{\min} , as the high degree cutoff can be determined in a similar fashion. The reader is advised to consult the discussion on systematic problems provided at the end of this section before implementing this procedure.

Fitting Procedure

As the degree distribution is typically provided as a list of positive integers k_{\min} , ..., k_{\max} , we aim to estimate γ from a discrete set of data points [47]. We use the citation network to illustrate the procedure. The network consists of N=384,362 nodes, each node representing a research paper published between 1890 and 2009 in journals published by the American Physical Society. The network has L = 2,353,984 links, each representing a citation from a published research paper to some other publication in the dataset (outside citations are ignored). For no particular reason, this is not the citation dataset listed in Table 4.1. See [48] for an overall characterization of this data. The steps of the fitting process are [47]:

1. Choose a value of K_{\min} between k_{\min} and k_{\max} . Estimate the value of the degree exponent corresponding to this K_{\min} using

$$\gamma = 1 + N \left[\sum_{i=1}^{N} \ln \frac{k_i}{K_{\min} - \frac{1}{2}} \right]^{-1}.$$
 (4.41)

Online Resource 4.2 Fitting power-law

The algorithmic tools to perform the fitting procedure described in this section are available at http://tuvalu.santafe.edu/~aaronc/powerlaws/.



2. With the obtained (γ , K_{\min}) parameter pair assume that the degree distribution has the form

$$p_k = \frac{1}{\zeta(\gamma, K_{\min})} k^{-\gamma}, \tag{4.42}$$

hence the associated cumulative distribution function (CDF) is

$$P_{k} = 1 - \frac{\zeta(\gamma, k)}{\zeta(\gamma, K_{\min})}.$$
(4.43)

3. Use the Kormogorov-Smirnov test to determine the maximum distance D between the CDF of the data S(k) and the fitted model provided by (4.43) with the selected (γ, k_{\min}) parameter pair,

$$D = \max_{k \ge K_{min}} |S(k) - P_k|. \tag{4.44}$$

Equation (4.44) identifies the degree for which the difference D between the empirical distribution S(k) and the fitted distribution (4.43) is the largest.

4. Repeat steps (1-3) by scanning the whole K_{\min} range from k_{\min} to k_{\max} . We aim to identify the K_{\min} value for which D provided by (4.44) is minimal. To illustrate the procedure, we plot D as a function of K_{\min} for the citation network (Figure 4.24b). The plot indicates that D is minimal for K_{\min} = 49, and the corresponding γ estimated by (4.41), representing the optimal fit, is γ =2.79. The standard error for the obtained degree exponent is

$$\sigma_{\gamma} = \frac{1}{\sqrt{N \left[\frac{\zeta''(\gamma, K_{\min})}{\zeta(\gamma, K_{\min})} - \left(\frac{\zeta'(\gamma, K_{\min})}{\zeta(\gamma, K_{\min})} \right)^{2} \right]}}$$
(4.45)

which implies that the best fit is $\gamma \pm \sigma_{\gamma}$. For the citation network we obtain σ_{γ} =0.003, hence γ =2.79(3).

Note that in order to estimate γ datasets smaller than N=50 should be treated with caution.

Goodness-of-fit

Just because we obtained a (γ, K_{\min}) pair that represents an optimal fit to our dataset, does not mean that the power law itself is a good model for the studied distribution. We therefore need to use a goodness-of-fit test, which generates a p-value that quantifies the plausibility of the power law hypothesis. The most often used procedure consists of the following steps:

1. Use the cumulative distribution (4.43) to estimate the KS distance between the real data and the best fit, that we denote by $D^{\rm real}$. This is step 3 above, taking the value of D for $K_{\rm min}$ that offered the best fit to the data. For the citation data we obtain $D^{\rm real} = 0.01158$ for $K_{\rm min} = 49$ (Figure 4.24c).

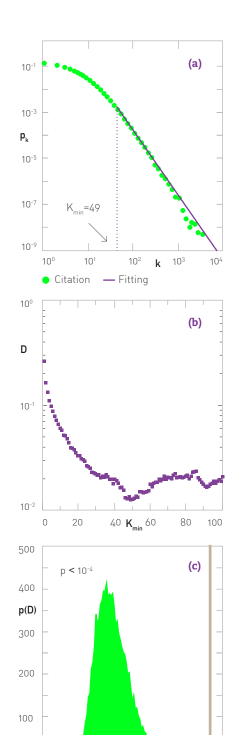


Figure 4.24
Maximum Likelihood Estimation

0.000

(a) The degree distribution p_k of the citation network, where the straight purple line represents the best fit based on the model (4.39).

0.005

0.010

- (b) The values of Kormogorov-Smirnov test vs. K_{\min} for the citation network.
- (c) p(D^{synthetic}) for M=10,000 synthetic datasets, where the grey line corresponds to the D^{real} value extracted for the citation network.

- 2. Use (4.42) to generate a degree sequence of N degrees (i.e. the same number of random numbers as the number of nodes in the original dataset) and substitute the obtained degree sequence for the empirical data, determining $D^{\rm synthetic}$ for this hypothetical degree sequence. Hence $D^{\rm synthetic}$ represents the distance between a synthetically generated degree sequence, consistent with our degree distribution, and the real data.
- 3. The goal is to see if the obtained $D^{\rm synthetic}$ is comparable to $D^{\rm real}$. For this we repeat step (2) M times ($M\gg 1$), and each time we generate a new degree sequence and determine the corresponding $D^{\rm synthetic}$, eventually obtaining the $p(D^{\rm synthetic})$ distribution. Plot $p(D^{\rm synthetic})$ and show as a vertical bar $D^{\rm real}$ (Figure 4.24c). If $D^{\rm real}$ is within the $p(D^{\rm synthetic})$ distribution, it means that the distance between the model providing the best fit and the empirical data is comparable with the distance expected from random degree samples chosen from the best fit distribution. Hence the power law is a reasonable model for the data. If, however, $D^{\rm real}$ falls outside the $p(D^{\rm synthetic})$ distribution, then the power law is not a good model some other function is expected to describe the original p_{ν} better.

While the distribution shown in Figure 4.24c may be in some cases useful to illustrate the statistical significance of the fit, in general it is better to assign a p-number to the fit, given by

$$p = \int_{D}^{\infty} P(D^{\text{synthetic}}) dD^{\text{synthetic}}.$$
 (4.46)

The closer p is to 1, the more likely that the difference between the empirical data and the model can be attributed to statistical fluctuations alone. If p is very small, the model is not a plausible fit to the data.

Typically, the model is accepted if p > 1%. For the citation network we obtain $p < 10^{-4}$, indicating that a pure power law is not a suitable model for the original degree distribution. This outcome is somewhat surprising, as the power-law nature of citation data has been documented repeatedly since 1960s [7, 8]. This failure indicates the limitation of the blind fitting to a power law, without an analytical understanding of the underlying distribution.

Fitting Real Distributions

To correct the problem, we note that the fitting model (4.44) eliminates all the data points with $k < K_{\min}$. As the citation network is fat tailed, choosing $K_{\min} = 49$ forces us to discard over 96% of the data points. Yet, there is statistically useful information in the $k < K_{\min}$ regime, that is ignored by the previous fit. We must introduce an alternate model that resolves this problem.

As we discussed in BOX 4.10, the degree distribution of many real networks, like the citation network, does not follow a pure power law. It often has low degree saturations and high degree cutoffs, described by

the form

$$p_{k} = \frac{1}{\sum_{k=1}^{L} (k' + k_{\text{sat}})^{-\gamma} e^{-k'/k_{\text{cut}}}} (k + k_{\text{sat}})^{-\gamma} e^{-k/k_{\text{cut}}}$$
(4.47)

and the associated CDF is

$$P_{k} = \frac{1}{\sum_{k'=1}^{k'=1} (k' + k_{\text{sat}})^{-\gamma} e^{-k'/k_{\text{cut}}}} \sum_{k'=1}^{k} (k' + k_{\text{sat}})^{-\gamma} e^{-k'/k_{\text{cut}}},$$
(4.48)

where k_{sat} and k_{cut} correspond to low-k saturation and the large-k cutoff, respectively. The difference between our earlier procedure and (4.47) is that we now do not discard the points that deviate from a pure power law, but instead use a function that offers a better fit to the whole degree distribution, from k_{min} to k_{max} .

Our goal is to find the fitting parameters $k_{\rm sat}$, $k_{\rm cut}$, and γ of the model (4.47), which we achieve through the following steps (Figure 4.25):

1. Pick a value for $k_{\rm sat}$ and $k_{\rm cut}$ between $K_{\rm min}$ and $K_{\rm max}$. Estimate the value of the degree exponent γ using the steepest descend method that maximizes the log-likelihood function

$$\log \mathcal{L}(\gamma \mid k_{\text{sat}}, k_{\text{cut}}) = \sum_{i=1}^{N} \log p(k_i \mid \gamma, k_{\text{sat}}, k_{\text{cut}}). \tag{4.49}$$

That is, for fixed $(k_{\text{sat}}, k_{\text{cut}})$ we vary γ until we find the maximum of (4.49).

- 2. With the obtained $\gamma(k_{\rm sat}, k_{\rm cut})$ assume that the degree distribution has the form (4.47). Calculate the Kormogorov Smirnov parameter D between the cumulative degree distribution (CDF) of the original data and the fitted model provided by (4.47).
- 3. Change $k_{\rm sat}$ and $k_{\rm cut}$, and repeat steps (1-3), scanning with $k_{\rm sat}$ from $k_{\rm min}$ = 0 to $k_{\rm max}$ and scanning with $k_{\rm cut}$ from $k_{\rm min}$ = k_0 to $k_{\rm max}$. The goal is to identify $k_{\rm sat}$ and $k_{\rm cut}$ values for which D is minimal. We illustrate this by plotting D in function of $k_{\rm sat}$ for several $k_{\rm cut}$ values in Figure 4.25a for our citation network. The ($k_{\rm sat}$, $k_{\rm cut}$) for which D is minimal, and the corresponding γ is provided by (4.41), represent the optimal parameters of the fit. For our dataset the optimal fit is obtained for $k_{\rm sat}$ = 12 and $k_{\rm cut}$ = 5,691, providing the degree exponent γ = 3.028. We find that now D for the real data is within the generated p distribution (Figure 4.25c), and the associated p-value is 69%.

Systematic Fitting Issues

The procedure described above may offer the impression that determining the degree exponent is a cumbersome but straightforward process. In reality these fitting methods have some well known limitations:

1. A pure power law is an idealized distribution that emerges in its

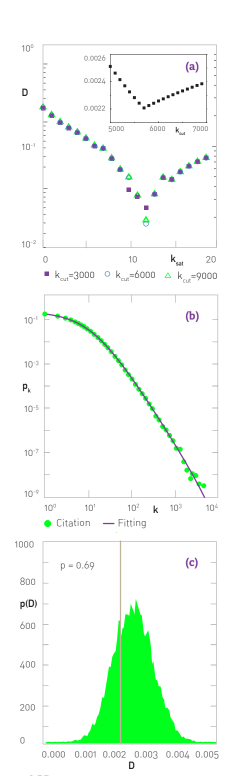


Figure 4.25
Estimating the Scaling Parameters for Citation
Networks

- (a) The Kormogorov-Smirnov parameter D vs. $k_{\rm sat}$ for $k_{\rm cut}$ = 3,000, 6,000, 9,000, respectively. The curve indicates that $k_{\rm sat}$ = 12 corresponds to the minimal D. Inset: D vs. $k_{\rm cut}$ for $k_{\rm sat}$ = 12, indicating that $k_{\rm cut}$ =5,691 minimizes D.
- (b) Degree distribution p_k where the straight line represents the best estimate from (a). Now the fit accurately captures the whole curve, not only its tail, or it did in **Figure** 4 24a
- (c) $p(D^{\text{synthetic}})$ for M=10,000 synthetic datasets. The grey line corresponds to the D^{real} value from the citation network.

form (4.1) only in simple models (CHAPTER 5). In reality, a whole range of processes contribute to the topology of real networks, affecting the precise shape of the degree distribution. These processes will be discussed in CHAPTER 6. If p_k does not follow a pure power law, the methods described above, designed to fit a power law to the data, will inevitably fail to detect statistical significance. While this finding can mean that the network is not scale-free, it most often means that we have not yet gained a proper understanding of the precise form of the degree distribution. Hence we are fitting the wrong functional form of p_k to the dataset.

2. The statistical tools used above to test the goodness-of-fit rely on the Kolmogorov-Smirnov criteria, which measures the maximum distance between the fitted model and the dataset. If almost all data points follow a perfect power law, but a *single* point for some reason deviates from the curve, we will loose the fit's statistical significance. In real systems there are numerous reasons for such local deviations that have little impact on the system's overall behavior. Yet, removing these "outliers" could be seen as data manipulation; if kept, however, one cannot detect the statistical significance of the power law fit.

A good example is provided by the actor network, whose degree distribution follows a power law for most degrees. There is, however, a prominent outlier at k=1,287, thanks to the 1956 movie *Around the World in Eighty Days*. This is the only movie where imdb.com the source of the actor network, lists all the normally uncredited extras in the cast. Hence the movie appears to have 1,288 actors. The second largest movie in the dataset has only 340 actors. Since each extra has links only to the 1,287 extras that played in the same movie, we have a local peak in p_k at k=1,287. Thanks to this peak, the degree distribution, fitted to a power law, fails to pass the Kolmogorov-Smirnov criteria. Indeed, as indicated in Table 4.3, neither the pure power law fit, nor a power law with high-degree cutoff offers a statistically significant fit. Yet, ultimately this single point does not alter the power law nature of the degreee distribution.

4. As a result of the issues discussed above, the methodology described to fit a power law distribution often predicts a small scaling regime, forcing us to remove a huge fraction of the nodes (often as many as

	λ	$k_{ m min}$	P-VALUE	PERCENTAGE	
Power Grid	0.517	4	0.91	12%	

99%, see Table 4.4) to obtain a statistically significant fit. Once plotted next to the original dataset, the obtained fit can be at times ridiculous, even if the method predits statistical significance.

Table 4.3
Exponential Fitting

For the power grid a power law degree distribution does not offer a statistically significant fit. Indeed, we will encounter numerous evidence that the underlying network is not scale-free. We used the fitting procedure described in this section to fit the exponential function $e^{-\lambda k}$ to the degree distribution of the power grid, obtaining a statistically significant fit. The table shows the obtained λ parameters, the k_{\min} over which the fit is valid, the obtained p-value, and the percentage of data points included in the fit.

In summary, estimating the degree exponent is still not yet an exact science. We continue to lack methods that would estimate the statistical significance in a manner that would be acceptable to a practitioner. The blind application of the tools describe above often leads to either fits that obviously do not capture the trends in the data, or to a false rejection of the power-law hypothesis. An important improvement is our ability to derive the expected form of the degree distribution, a problem discussed in CHAP-

TER 6.

K ^{-γ} ;[K _{min} ,∞]			$\left(k + k_{sat}\right)^{-\gamma} e^{-k/k_{cut}}$					
	γ	K_{\min}	P-VALUE	PERCENT	γ	k _{sat}	k _{cut}	P-VALUE
INTERNET	3.42	72	0.13	0.6%	3.55	8	8500	0.00
www (IN)	2.00	1	0.00	100%	1.97	0	660	0.00
www (OUT)	2.31	7	0.00	15%	2.82	8	8500	0.00
POWER GRID	4.00	5	0.00	12%	8.56	19	14	0.00
MOBILE PHONE CALLS (IN)	4.69	9	0.34	2.6%	6.95	15	10	0.00
MOBILE PHONE CALLS [OUT]	5.01	11	0.77	1.7%	7.23	15	10	0.00
EMAIL-PRE (IN)	3.43	88	0.11	0.2%	2.27	0	8500	0.00
EMAIL-PRE (OUT)	2.03	3	0.00	1.2%	2.55	0	8500	0.00
SCIENCE COLLABORATION	3.35	25	0.0001	5.4%	1.50	17	12	0.00
ACTOR NETWORK	2.12	54	0.00	33%	-	-	-	0.00
CITATION NETWORK (IN)	2.79	51	0.00	3.0%	3.03	12	5691	0.69
CITATION NETWORK (OUT)	4.00	19	0.00	14%	-0.16	5	10	0.00
E.COLI METABOLISM	2.43	3	0.00	57%	3.85	19	12	0.00
E.COLI METABOLISM (OUT)	2.90	5	0.00	34%	2.56	15	10	0.00
YEAST PROTEIN INTERACTIONS	2.89	7	0.67	8.3%	2.95	2	90	0.52

Table 4.4

Fitting Parameters for Real Networks

The estimated degree exponents and the appropriate fit parameters for the reference networks studied in this book. We implement two fitting strategies, the first aiming to fit a pure power law in the region (K_{\min}, ∞) and the second fits a power law with saturation and exponential cutoff to the whole dataset. In the table we show the obtained γ exponent and K_{\min} for the fit with the best statistical significance, the *p*-value for the best fit and the percentage of the data included in the fit. In the second case we again show the exponent γ , the two fit parameters, k_{sat} and k_{cut} , and the p-value of the obtained fit. Note that p > 0.01is considered to be statistically significant.

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