Foundations of Data Science*

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1 Introduction

Computer science as an academic discipline began in the 1960's. Emphasis was on programming languages, compilers, operating systems, and the mathematical theory that supported these areas. Courses in theoretical computer science covered finite automata, regular expressions, context free languages, and computability. In the 1970's, the study of algorithms was added as an important component of theory. The emphasis was on making computers useful. Today, a fundamental change is taking place and the focus is more on applications. There are many reasons for this change. The merging of computing and communications has played an important role. The enhanced ability to observe, collect and store data in the natural sciences, in commerce, and in other fields calls for a change in our understanding of data and how to handle it in the modern setting. The emergence of the web and social networks as central aspects of daily life presents both opportunities and challenges for theory.

While traditional areas of computer science remain highly important, increasingly researchers of the future will be involved with using computers to understand and extract usable information from massive data arising in applications, not just how to make computers useful on specific well-defined problems. With this in mind we have written this book to cover the theory likely to be useful in the next 40 years, just as an understanding of automata theory, algorithms and related topics gave students an advantage in the last 40 years. One of the major changes is the switch from discrete mathematics to more of an emphasis on probability, statistics, and numerical methods.

Early drafts of the book have been used for both undergraduate and graduate courses. Background material needed for an undergraduate course has been put in the appendix. For this reason, the appendix has homework problems.

This book starts with the treatment of high dimensional geometry. Modern data in diverse fields such as Information Processing, Search, Machine Learning, etc., is often represented advantageously as vectors with a large number of components. This is so even in cases when the vector representation is not the natural first choice. Our intuition from two or three dimensional space can be surprisingly off the mark when it comes to high dimensional space. Chapter 2 works out the fundamentals needed to understand the differences. The emphasis of the chapter, as well as the book in general, is to get across the mathematical foundations rather than dwell on particular applications that are only briefly described.

The mathematical areas most relevant to dealing with high-dimensional data are matrix algebra and algorithms. We focus on singular value decomposition, a central tool in this area. Chapter 3 gives a from-first-principles description of this. Applications of singular value decomposition include principal component analysis, a widely used technique which we touch upon, as well as modern applications to statistical mixtures of probability densities, discrete optimization, etc., which are described in more detail.

Central to our understanding of large structures, like the web and social networks, is building models to capture essential properties of these structures. The simplest model is that of a random graph formulated by Erdös and Renyi, which we study in detail proving that certain global phenomena, like a giant connected component, arise in such structures with only local choices. We also describe other models of random graphs.

One of the surprises of computer science over the last two decades is that some domainindependent methods have been immensely successful in tackling problems from diverse areas. Machine learning is a striking example. We describe the foundations of machine learning, both algorithms for optimizing over given training examples, as well as the theory for understanding when such optimization can be expected to lead to good performance on new, unseen data, including important measures such as Vapnik-Chervonenkis dimension. Another important domain-independent technique is based on Markov chains. The underlying mathematical theory, as well as the connections to electrical networks, forms the core of our chapter on Markov chains.

The field of algorithms has traditionally assumed that the input data to a problem is presented in random access memory, which the algorithm can repeatedly access. This is not feasible for modern problems. The streaming model and other models have been formulated to better reflect this. In this setting, sampling plays a crucial role and, indeed, we have to sample on the fly. In Chapter 7 we study how to draw good samples efficiently and how to estimate statistical and linear algebra quantities, with such samples.

Another important tool for understanding data is clustering, dividing data into groups of similar objects. After describing some of the basic methods for clustering, such as the k-means algorithm, we focus on modern developments in understanding these, as well as newer algorithms. The chapter ends with a study of clustering criteria.

This book also covers graphical models and belief propagation, ranking and voting, sparse vectors, and compressed sensing. The appendix includes a wealth of background material.

A word about notation in the book. To help the student, we have adopted certain notations, and with a few exceptions, adhered to them. We use lower case letters for scalar variables and functions, bold face lower case for vectors, and upper case letters for matrices. Lower case near the beginning of the alphabet tend to be constants, in the middle of the alphabet, such as i, j, and k, are indices in summations, n and m for integer sizes, and x, y and z for variables. If A is a matrix its elements are a_{ij} and its rows are \mathbf{a}_i . If \mathbf{a}_i is a vector its coordinates are a_{ij} . Where the literature traditionally uses a symbol for a quantity, we also used that symbol, even if it meant abandoning our convention. If we have a set of points in some vector space, and work with a subspace, we use n for the

number of points, d for the dimension of the space, and k for the dimension of the subspace.

The term "almost surely" means with probability one. We use $\ln n$ for the natural logarithm and $\log n$ for the base two logarithm. If we want base ten, we will use \log_{10} . To simplify notation and to make it easier to read we use $E^2(1-x)$ for $(E(1-x))^2$ and $E(1-x)^2$ for $E((1-x)^2)$. When we say "randomly select" some number of points from a given probability distribution D, independence is always assumed unless otherwise stated.

2 High-Dimensional Space

2.1 Introduction

High dimensional data has become very important. However, high dimensional space is very different from the two and three dimensional spaces we are familiar with. Generate n points at random in d-dimensions where each coordinate is a zero mean, unit variance Gaussian. For sufficiently large d, with high probability the distances between all pairs of points will be essentially the same. Also the volume of the unit ball in d-dimensions, the set of all points \mathbf{x} such that $|\mathbf{x}| \leq 1$, goes to zero as the dimension goes to infinity. The volume of a high dimensional unit ball is concentrated near its surface and is also concentrated at its equator. These properties have important consequences which we will consider.

2.2 The Law of Large Numbers

If one generates random points in *d*-dimensional space using a Gaussian to generate coordinates, the distance between all pairs of points will be essentially the same when *d* is large. The reason is that the square of the distance between two points \mathbf{y} and \mathbf{z} ,

$$|\mathbf{y} - \mathbf{z}|^2 = \sum_{i=1}^d (y_i - z_i)^2,$$

is the sum of d independent random variables. If one averages n independent samples of a random variable x of bounded variance, the result will be close to the expected value of x. In the above summation there are d samples where each sample is the squared distance in a coordinate between the two points \mathbf{y} and \mathbf{z} . Here we give a general bound called the Law of Large Numbers. Specifically, the Law of Large Numbers states that

$$\operatorname{Prob}\left(\left|\frac{x_1 + x_2 + \dots + x_n}{n} - E(x)\right| \ge \epsilon\right) \le \frac{\operatorname{Var}(x)}{n\epsilon^2}.$$
(2.1)

The larger the variance of the random variable, the greater the probability that the error will exceed ϵ . Thus the variance of x is in the numerator. The number of samples n is in the denominator since the more values that are averaged, the smaller the probability that the difference will exceed ϵ . Similarly the larger ϵ is, the smaller the probability that the difference will exceed ϵ and hence ϵ is in the denominator. Notice that squaring ϵ makes the fraction a dimensionless quantity.

We use two inequalities to prove the Law of Large Numbers. The first is Markov's inequality which states that the probability that a nonnegative random variable exceeds a is bounded by the expected value of the variable divided by a.

Theorem 2.1 (Markov's inequality) Let x be a nonnegative random variable. Then for a > 0,

$$Prob(x \ge a) \le \frac{E(x)}{a}.$$

Proof: For a continuous nonnegative random variable x with probability density p,

$$E(x) = \int_{0}^{\infty} xp(x)dx = \int_{0}^{a} xp(x)dx + \int_{a}^{\infty} xp(x)dx$$
$$\geq \int_{a}^{\infty} xp(x)dx \ge a \int_{a}^{\infty} p(x)dx = a \operatorname{Prob}(x \ge a)$$

Thus, $\operatorname{Prob}(x \ge a) \le \frac{E(x)}{a}$.

The same proof works for discrete random variables with sums instead of integrals.

Corollary 2.2 $Prob(x \ge bE(x)) \le \frac{1}{b}$

Markov's inequality bounds the tail of a distribution using only information about the mean. A tighter bound can be obtained by also using the variance of the random variable.

Theorem 2.3 (Chebyshev's inequality) Let x be a random variable. Then for c > 0,

$$Prob\left(|x - E(x)| \ge c\right) \le \frac{Var(x)}{c^2}$$

Proof: Prob $(|x - E(x)| \ge c) = \operatorname{Prob}(|x - E(x)|^2 \ge c^2)$. Let $y = |x - E(x)|^2$. Note that y is a nonnegative random variable and E(y) = Var(x), so Markov's inequality can be applied giving:

$$\operatorname{Prob}(|x - E(x)| \ge c) = \operatorname{Prob}\left(|x - E(x)|^2 \ge c^2\right) \le \frac{E(|x - E(x)|^2)}{c^2} = \frac{Var(x)}{c^2}.$$

The Law of Large Numbers follows from Chebyshev's inequality together with facts about independent random variables. Recall that:

$$E(x + y) = E(x) + E(y),$$

$$Var(x - c) = Var(x),$$

$$Var(cx) = c^{2}Var(x).$$

Also, if x and y are independent, then E(xy) = E(x)E(y). These facts imply that if x and y are independent then Var(x+y) = Var(x) + Var(y), which is seen as follows:

$$Var(x+y) = E(x+y)^2 - E^2(x+y)$$

= $E(x^2 + 2xy + y^2) - (E^2(x) + 2E(x)E(y) + E^2(y))$
= $E(x^2) - E^2(x) + E(y^2) - E^2(y) = Var(x) + Var(y),$

where we used independence to replace E(2xy) with 2E(x)E(y).

Theorem 2.4 (Law of large numbers) Let x_1, x_2, \ldots, x_n be n independent samples of a random variable x. Then

$$Prob\left(\left|\frac{x_1+x_2+\dots+x_n}{n}-E(x)\right| \ge \epsilon\right) \le \frac{Var(x)}{n\epsilon^2}$$

Proof: By Chebychev's inequality

$$\operatorname{Prob}\left(\left|\frac{x_1 + x_2 + \dots + x_n}{n} - E(x)\right| \ge \epsilon\right) \le \frac{\operatorname{Var}\left(\frac{x_1 + x_2 + \dots + x_n}{n}\right)}{\epsilon^2}$$
$$= \frac{1}{n^2 \epsilon^2} \operatorname{Var}(x_1 + x_2 + \dots + x_n)$$
$$= \frac{1}{n^2 \epsilon^2} \left(\operatorname{Var}(x_1) + \operatorname{Var}(x_2) + \dots + \operatorname{Var}(x_n)\right)$$
$$= \frac{\operatorname{Var}(x)}{n \epsilon^2}.$$

The Law of Large Numbers is quite general, applying to any random variable x of finite variance. Later we will look at tighter concentration bounds for spherical Gaussians and sums of 0-1 valued random variables.

As an application of the Law of Large Numbers, let \mathbf{z} be a *d*-dimensional random point whose coordinates are each selected from a zero mean, $\frac{1}{2\pi}$ variance Gaussian. We set the variance to $\frac{1}{2\pi}$ so the Gaussian probability density equals one at the origin and is bounded below throughout the unit ball by a constant. By the Law of Large Numbers, the square of the distance of \mathbf{z} to the origin will be $\Theta(d)$ with high probability. In particular, there is vanishingly small probability that such a random point \mathbf{z} would lie in the unit ball. This implies that the integral of the probability density over the unit ball must be vanishingly small. On the other hand, the probability density in the unit ball is bounded below by a constant. We thus conclude that the unit ball must have vanishingly small volume.

Similarly if we draw two points \mathbf{y} and \mathbf{z} from a *d*-dimensional Gaussian with unit variance in each direction, then $|\mathbf{y}|^2 \approx d$, $|\mathbf{z}|^2 \approx d$, and $|\mathbf{y} - \mathbf{z}|^2 \approx 2d$ (since $E(y_i - z_i)^2 = E(y_i^2) + E(z_i^2) - 2E(y_i z_i) = 2$ for all *i*.) Thus by the Pythagorean theorem, the random

d-dimensional \mathbf{y} and \mathbf{z} must be approximately orthogonal. This implies that if we scale these random points to be unit length and call \mathbf{y} the North Pole, much of the surface area of the unit ball must lie near the equator. We will formalize these and related arguments in subsequent sections.

We now state a general theorem on probability tail bounds for a sum of independent random variables. Tail bounds for sums of Bernoulli, squared Gaussian and Power Law distributed random variables can all be derived from this. The table below summarizes some of the results.

Theorem 2.5 (Master Tail Bounds Theorem) Let $x = x_1 + x_2 + \cdots + x_n$, where x_1, x_2, \ldots, x_n are mutually independent random variables with zero mean and variance at most σ^2 . Let $0 \le a \le \sqrt{2}n\sigma^2$. Assume that $|E(x_i^s)| \le \sigma^2 s!$ for $s = 3, 4, \ldots, \lfloor (a^2/4n\sigma^2) \rfloor$. Then,

$$Prob(|x| \ge a) \le 3e^{-a^2/(12n\sigma^2)}.$$

The elementary proof of Theorem 12.5 is given in the appendix. For a brief intuition, consider applying Markov's inequality to the random variable x^r where r is a large even number. Since r is even, x^r is non-negative, and thus $\operatorname{Prob}(|x| \ge a) = \operatorname{Prob}(x^r \ge a^r) \le E(x^r)/a^r$. If $E(x^r)$ is not too large, we will get a good bound. To compute $E(x^r)$, write E(x) as $E(x_1 + \ldots + x_n)^r$ and distribute the polynomial into its terms. Use the fact that by independence $E(x_i^{r_i}x_j^{r_j}) = E(x_i^{r_i})E(x_j^{r_j})$ to get a collection of simpler expectations that can be bounded using our assumption that $|E(x_i^s)| \le \sigma^2 s!$. For the full proof, see the appendix.

	Condition	Tail bound	Notes
Markov	$x \ge 0$	$\Pr(x \ge a) \le \frac{E(x)}{a}$	
Chebychev	Any x	$\Pr(x - E(x) \ge a) \le \frac{\operatorname{Var}(x)}{a^2}$	
Chernoff	$x = x_1 + x_2 + \dots + x_n$	$\Pr(x - E(x) \ge \varepsilon E(x)) \le$	From Thm 12.5
	x_i i.i.d. Bernoulli; $\varepsilon \in [0, 1]$	$3\exp(-c\varepsilon^2 E(x))$	Appendix
Higher Moments	r pos. even int.	$\Pr(x \ge a) \le E(x^r)/a^r$	Markov on x^r
Gaussian	$x = \sqrt{x_1^2 + x_2^2 + \dots + x_n^2}$	$\Pr(x - \sqrt{n} \ge \beta) \le$	From Thm 12.5.
Annulus	$x_i \sim N(0,1); \beta \leq \sqrt{n}$ indep.	$3\exp(-c\beta^2)$	Section (2.6)
Power Law	$x = x_1 + x_2 + \ldots + x_n; x_i \text{ i.i.d}$	$\Pr(x - E(x) \ge \varepsilon E(x)) \le$	From Thm 12.5
for x_i ; order $k \ge 4$	$\varepsilon \leq 1/k^2$	$\leq (4/\varepsilon^2 kn)^{(k-3)/2}$	Appendix

Table of Tail Bounds

2.3 The Geometry of High Dimensions

An important property of high-dimensional objects is that most of their volume is near the surface. Consider any object A in \mathbb{R}^d . Now shrink A by a small amount ϵ to produce a new object $(1 - \epsilon)A = \{(1 - \epsilon)x | x \in A\}$. Then volume $((1 - \epsilon)A) = (1 - \epsilon)^d$ volume(A). To see that this is true, partition A into infinitesimal cubes. Then, $(1 - \varepsilon)A$ is the union of a set of cubes obtained by shrinking the cubes in A by a factor of $1 - \varepsilon$. Now, when

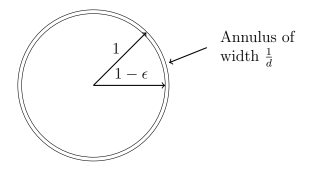


Figure 2.1: Most of the volume of the *d*-dimensional ball of radius r is contained in an annulus of width O(r/d) near the boundary.

we shrink each of the d sides of a d-dimensional cube by a factor f, its volume shrinks by a factor of f^d . Using the fact that $1 - x \le e^{-x}$, for any object A in \mathbb{R}^d we have:

$$\frac{\operatorname{volume}((1-\epsilon)A)}{\operatorname{volume}(A)} = (1-\epsilon)^d \leq e^{-\epsilon d}$$

Fixing ϵ and letting $d \to \infty$, the above quantity rapidly approaches zero. This means that nearly all of the volume of A must be in the portion of A that does not belong to the region $(1 - \epsilon)A$.

Let S denote the unit ball in d dimensions, that is, the set of points within distance one of the origin. An immediate implication of the above observation is that at least a $1 - e^{-\epsilon d}$ fraction of the volume of the unit ball is concentrated in $S \setminus (1 - \epsilon)S$, namely in a small annulus of width ϵ at the boundary. In particular, most of the volume of the d-dimensional unit ball is contained in an annulus of width O(1/d) near the boundary. If the ball is of radius r, then the annulus width is $O\left(\frac{r}{d}\right)$.

2.4 Properties of the Unit Ball

We now focus more specifically on properties of the unit ball in *d*-dimensional space. We just saw that most of its volume is concentrated in a small annulus of width O(1/d) near the boundary. Next we will show that in the limit as *d* goes to infinity, the volume of the ball goes to zero. This result can be proven in several ways. Here we use integration.

2.4.1 Volume of the Unit Ball

To calculate the volume V(d) of the unit ball in \mathbb{R}^d , one can integrate in either Cartesian or polar coordinates. In Cartesian coordinates the volume is given by

$$V(d) = \int_{x_1=-1}^{x_1=1} \int_{x_2=-\sqrt{1-x_1^2}}^{x_2=\sqrt{1-x_1^2}} \cdots \int_{x_d=-\sqrt{1-x_1^2-\cdots-x_{d-1}^2}}^{x_d=\sqrt{1-x_1^2-\cdots-x_{d-1}^2}} dx_d \cdots dx_2 dx_1.$$

Since the limits of the integrals are complicated, it is easier to integrate using polar coordinates. In polar coordinates, V(d) is given by

$$V(d) = \int_{S^d} \int_{r=0}^1 r^{d-1} dr d\Omega.$$

Since the variables Ω and r do not interact,

$$V(d) = \int_{S^d} d\Omega \int_{r=0}^1 r^{d-1} dr = \frac{1}{d} \int_{S^d} d\Omega = \frac{A(d)}{d}$$

where A(d) is the surface area of the *d*-dimensional unit ball. For instance, for d = 3 the surface area is 4π and the volume is $\frac{4}{3}\pi$. The question remains, how to determine the surface area $A(d) = \int_{S^d} d\Omega$ for general d.

Consider a different integral

$$I(d) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \cdots \int_{-\infty}^{\infty} e^{-(x_1^2 + x_2^2 + \cdots + x_d^2)} dx_d \cdots dx_2 dx_1.$$

Including the exponential allows integration to infinity rather than stopping at the surface of the sphere. Thus, I(d) can be computed by integrating in both Cartesian and polar coordinates. Integrating in polar coordinates will relate I(d) to the surface area A(d). Equating the two results for I(d) allows one to solve for A(d).

First, calculate I(d) by integration in Cartesian coordinates.

$$I(d) = \left[\int_{-\infty}^{\infty} e^{-x^2} dx\right]^d = \left(\sqrt{\pi}\right)^d = \pi^{\frac{d}{2}}.$$

Here, we have used the fact that $\int_{-\infty}^{\infty} e^{-x^2} dx = \sqrt{\pi}$. For a proof of this, see Section ?? of the appendix. Next, calculate I(d) by integrating in polar coordinates. The volume of the differential element is $r^{d-1}d\Omega dr$. Thus,

$$I(d) = \int_{S^d} d\Omega \int_0^\infty e^{-r^2} r^{d-1} dr.$$

The integral $\int_{S^d} d\Omega$ is the integral over the entire solid angle and gives the surface area, A(d), of a unit sphere. Thus, $I(d) = A(d) \int_{0}^{\infty} e^{-r^2} r^{d-1} dr$. Evaluating the remaining integral gives

$$\int_{0}^{\infty} e^{-r^{2}} r^{d-1} dr = \int_{0}^{\infty} e^{-t} t^{\frac{d-1}{2}} \left(\frac{1}{2} t^{-\frac{1}{2}} dt\right) = \frac{1}{2} \int_{0}^{\infty} e^{-t} t^{\frac{d}{2}} - \frac{1}{2} dt = \frac{1}{2} \Gamma\left(\frac{d}{2}\right)$$

and hence, $I(d) = A(d)\frac{1}{2}\Gamma\left(\frac{d}{2}\right)$ where the Gamma function $\Gamma(x)$ is a generalization of the factorial function for noninteger values of x. $\Gamma(x) = (x-1)\Gamma(x-1)$, $\Gamma(1) = \Gamma(2) = 1$, and $\Gamma\left(\frac{1}{2}\right) = \sqrt{\pi}$. For integer x, $\Gamma(x) = (x-1)!$.

Combining $I(d) = \pi^{\frac{d}{2}}$ with $I(d) = A(d) \frac{1}{2} \Gamma\left(\frac{d}{2}\right)$ yields

$$A(d) = \frac{\pi^2}{\frac{1}{2}\Gamma\left(\frac{d}{2}\right)}$$

establishing the following lemma.

Lemma 2.6 The surface area A(d) and the volume V(d) of a unit-radius ball in d dimensions are given by

$$A\left(d\right) = \frac{2\pi^{\frac{d}{2}}}{\Gamma\left(\frac{d}{2}\right)} \quad and \quad V\left(d\right) = \frac{2}{d} \frac{\pi^{\frac{d}{2}}}{\Gamma\left(\frac{d}{2}\right)}.$$

To check the formula for the volume of a unit ball, note that $V(2) = \pi$ and $V(3) = \frac{2}{3} \frac{\pi^{\frac{3}{2}}}{\Gamma(\frac{3}{2})} = \frac{4}{3}\pi$, which are the correct volumes for the unit balls in two and three dimensions. To check the formula for the surface area of a unit ball, note that $A(2) = 2\pi$ and $A(3) = \frac{2\pi^{\frac{3}{2}}}{\frac{1}{2}\sqrt{\pi}} = 4\pi$, which are the correct surface areas for the unit ball in two and three dimensions. Note that $\pi^{\frac{d}{2}}$ is an exponential in $\frac{d}{2}$ and $\Gamma(\frac{d}{2})$ grows as the factorial of $\frac{d}{2}$. This implies that $\lim_{d\to\infty} V(d) = 0$, as claimed.

2.4.2 Most of the Volume is Near the Equator

An interesting fact about the unit ball in high dimensions is that most of its volume is concentrated near its equator no matter what direction one uses to define the North Pole and hence the "equator". Arbitrarily letting x_1 denote "north", most of the volume of the unit ball has $|x_1| = O(1/\sqrt{d})$. Using this fact, we will show that two random points in the unit ball are with high probability nearly orthogonal, and also give an alternative proof from the one in Section 2.4.1 that the volume of the unit ball goes to zero as $d \to \infty$.

Theorem 2.7 For $c \ge 1$ and $d \ge 3$, at least a $1 - \frac{2}{c}e^{-c^2/2}$ fraction of the volume of the d-dimensional unit ball has $|x_1| \le \frac{c}{\sqrt{d-1}}$.

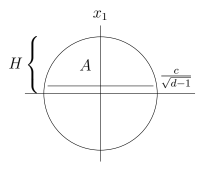


Figure 2.2: Most of the volume of the upper hemisphere of the *d*-dimensional ball is below the plane $x_1 = \frac{c}{\sqrt{d-1}}$.

Proof: By symmetry we just need to prove that at most a $\frac{2}{c}e^{-c^2/2}$ fraction of the half of the ball with $x_1 \ge 0$ has $x_1 \ge \frac{c}{\sqrt{d-1}}$. Let A denote the portion of the ball with $x_1 \ge \frac{c}{\sqrt{d-1}}$ and let H denote the upper hemisphere. We will then show that the ratio of the volume of A to the volume of H goes to zero by calculating an upper bound on volume(H) and a lower bound on volume(A) and proving that

$$\frac{\text{volume}(A)}{\text{volume}(H)} \le \frac{\text{upper bound volume}(A)}{\text{lower bound volume}(H)} = \frac{2}{c}e^{-\frac{c^2}{2}}.$$

To calculate the volume of A, integrate an incremental volume that is a disk of width dx_1 and whose face is a ball of dimension d-1 and radius $\sqrt{1-x_1^2}$. The surface area of the disk is $(1-x_1^2)^{\frac{d-1}{2}}V(d-1)$ and the volume above the slice is

volume(A) =
$$\int_{\frac{c}{\sqrt{d-1}}}^{1} (1 - x_1^2)^{\frac{d-1}{2}} V(d-1) dx_1$$

To get an upper bound on the above integral, use $1 - x \leq e^{-x}$ and integrate to infinity. To integrate, insert $\frac{x_1\sqrt{d-1}}{c}$, which is greater than one in the range of integration, into the integral. Then

Now

$$\int_{\frac{c}{\sqrt{d-1}}}^{\infty} x_1 e^{-\frac{d-1}{2}x_1^2} dx_1 = -\frac{1}{d-1} e^{-\frac{d-1}{2}x_1^2} \Big|_{\frac{c}{\sqrt{d-1}}}^{\infty} = \frac{1}{d-1} e^{-\frac{c^2}{2}}$$

Thus, an upper bound on volume(A) is $\frac{V(d-1)}{c\sqrt{d-1}}e^{-\frac{c^2}{2}}$.

The volume of the hemisphere below the plane $x_1 = \frac{1}{\sqrt{d-1}}$ is a lower bound on the entire volume of the upper hemisphere and this volume is at least that of a cylinder of height $\frac{1}{\sqrt{d-1}}$ and radius $\sqrt{1-\frac{1}{d-1}}$. The volume of the cylinder is $V(d-1)(1-\frac{1}{d-1})^{\frac{d-1}{2}}\frac{1}{\sqrt{d-1}}$. Using the fact that $(1-x)^a \ge 1 - ax$ for $a \ge 1$, the volume of the cylinder is at least $\frac{V(d-1)}{2\sqrt{d-1}}$ for $d \ge 3$.

Thus,

ratio
$$\leq \frac{\text{upper bound above plane}}{\text{lower bound total hemisphere}} = \frac{\frac{V(d-1)}{c\sqrt{d-1}}e^{-\frac{c^2}{2}}}{\frac{V(d-1)}{2\sqrt{d-1}}} = \frac{2}{c}e^{-\frac{c^2}{2}}$$

One might ask why we computed a lower bound on the total hemisphere since it is one half of the volume of the unit ball which we already know. The reason is that the volume of the upper hemisphere is $\frac{1}{2}V(d)$ and we need a formula with V(d-1) in it to cancel the V(d-1) in the numerator.

Near orthogonality. One immediate implication of the above analysis is that if we draw two points at random from the unit ball, with high probability their vectors will be nearly orthogonal to each other. Specifically, from our previous analysis in Section 2.3, with high probability both will have length 1 - O(1/d). From our analysis above, if we define the vector in the direction of the first point as "north", with high probability the second will have a projection of only $\pm O(1/\sqrt{d})$ in this direction. This implies that with high probability, the angle between the two vectors will be $\pi/2 \pm O(1/\sqrt{d})$. In particular, we have the theorem:

Theorem 2.8 Consider drawing n points $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_n}$ at random from the unit ball. With probability 1 - O(1/n)

1. $|\mathbf{x_i}| \ge 1 - \frac{2\ln n}{d}$ for all *i*, and 2. $|\mathbf{x_i} \cdot \mathbf{x_j}| \le \frac{\sqrt{6\ln n}}{\sqrt{d-1}}$ for all $i \ne j$.

Proof: For the first part, for any fixed *i*, by the analysis of Section 2.3 $\operatorname{Prob}(|\mathbf{x}_i| < 1 - \frac{2\ln n}{d}) \leq e^{-(\frac{2\ln n}{d})d} = 1/n^2$. So, by the union bound, the probability there exists *i* such that $|\mathbf{x}_i| < 1 - \frac{2\ln n}{d}$ is at most 1/n. For the second part, there are $\binom{n}{2}$ pairs *i* and *j*, and for each such pair, if we define \mathbf{x}_i as "north", the probability that the projection of \mathbf{x}_j onto the "North" direction is more than $\frac{\sqrt{6\ln n}}{\sqrt{d-1}}$ (a necessary condition for the dot-product to be large) is at most $O(e^{-\frac{6\ln n}{2}}) = O(n^{-3})$ by Theorem 2.7. Thus, this condition is violated with probability at most $O\left(\binom{n}{2}n^{-3}\right) = O(1/n)$ as well.

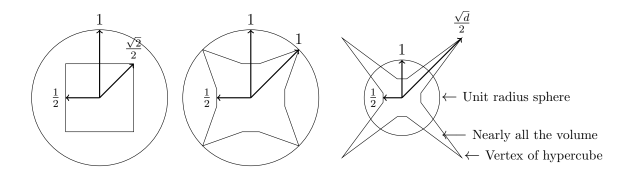


Figure 2.3: Illustration of the relationship between the sphere and the cube in 2, 4, and d-dimensions.

Alternative proof that volume goes to zero. Another immediate implication of Theorem 2.7 is that as $d \to \infty$, the volume of the ball approaches zero. Specifically, setting $c = 2\sqrt{\ln d}$ in Theorem 2.7, the fraction of the volume of the ball with $|x_1| \ge \frac{c}{\sqrt{d-1}}$ is at most:

$$\frac{2}{c}e^{-\frac{c^2}{2}} = \frac{1}{\sqrt{\ln d}}e^{-2\ln d} = \frac{1}{d^2\sqrt{\ln d}} < \frac{1}{d^2}.$$

Since this is true for each of the d dimensions, by a union bound at most a $O(\frac{1}{d}) \leq \frac{1}{2}$ fraction of the volume of the ball lies outside the cube of side-length $2\frac{c}{\sqrt{d-1}}$. Thus, the ball has volume at most twice that of this cube. This cube has volume $(\frac{16 \ln d}{d-1})^{d/2}$, and this quantity goes to zero as $d \to \infty$. Thus the volume of the ball goes to zero as well.

Discussion. One might wonder how it can be that nearly all the points in the unit ball are very close to the surface and yet at the same time nearly all points are in a box of side-length $O\left(\frac{\ln d}{d-1}\right)$. The answer is to remember that points on the surface of the ball satisfy $x_1^2 + x_2^2 + \ldots + x_d^2 = 1$, so for each coordinate *i*, a typical value will be $\pm O\left(\frac{1}{\sqrt{d}}\right)$. In fact, it is often helpful to think of picking a random point on the sphere as very similar to picking a random point of the form $\left(\pm \frac{1}{\sqrt{d}}, \pm \frac{1}{\sqrt{d}}, \ldots \pm \frac{1}{\sqrt{d}}\right)$.

2.5 Generating Points Uniformly at Random from a Ball

Consider generating points uniformly at random on the surface of the unit ball. For the 2-dimensional version of generating points on the circumference of a unit-radius circle, independently generate each coordinate uniformly at random from the interval [-1, 1]. This produces points distributed over a square that is large enough to completely contain the unit circle. Project each point onto the unit circle. The distribution is not uniform since more points fall on a line from the origin to a vertex of the square than fall on a line from the origin to the midpoint of an edge of the square due to the difference in length. To solve this problem, discard all points outside the unit circle and project the remaining points onto the circle.

In higher dimensions, this method does not work since the fraction of points that fall inside the ball drops to zero and all of the points would be thrown away. The solution is to generate a point each of whose coordinates is an independent Gaussian variable. Generate x_1, x_2, \ldots, x_d , using a zero mean, unit variance Gaussian, namely, $\frac{1}{\sqrt{2\pi}} \exp(-x^2/2)$ on the real line.¹ Thus, the probability density of **x** is

$$p(\mathbf{x}) = \frac{1}{(2\pi)^{\frac{d}{2}}} e^{-\frac{x_1^2 + x_2^2 + \dots + x_d^2}{2}}$$

and is spherically symmetric. Normalizing the vector $\mathbf{x} = (x_1, x_2, \dots, x_d)$ to a unit vector, namely $\frac{\mathbf{x}}{|\mathbf{x}|}$, gives a distribution that is uniform over the surface of the sphere. Note that once the vector is normalized, its coordinates are no longer statistically independent.

To generate a point \mathbf{y} uniformly over the ball (surface and interior), scale the point $\frac{\mathbf{x}}{|\mathbf{x}|}$ generated on the surface by a scalar $\rho \in [0, 1]$. What should the distribution of ρ be as a function of r? It is certainly not uniform, even in 2 dimensions. Indeed, the density of ρ at r is proportional to r for d = 2. For d = 3, it is proportional to r^2 . By similar reasoning, the density of ρ at distance r is proportional to r^{d-1} in d dimensions. Solving $\int_{r=0}^{r=1} cr^{d-1}dr = 1$ (the integral of density must equal 1) we should set c = d. Another way to see this formally is that the volume of the radius r ball in d dimensions is $r^d V(d)$. The density at radius r is exactly $\frac{d}{dr}(r^d V_d) = dr^{d-1}V_d$. So, pick $\rho(r)$ with density equal to dr^{d-1} for r over [0, 1].

We have succeeded in generating a point

$$\mathbf{y} = \rho \frac{\mathbf{x}}{|\mathbf{x}|}$$

uniformly at random from the unit ball by using the convenient spherical Gaussian distribution. In the next sections, we will analyze the spherical Gaussian in more detail.

2.6 Gaussians in High Dimension

A 1-dimensional Gaussian has its mass close to the origin. However, as the dimension is increased something different happens. The d-dimensional spherical Gaussian with zero

¹One might naturally ask: "how do you generate a random number from a 1-dimensional Gaussian?" To generate a number from any distribution given its cumulative distribution function P, first select a uniform random number $u \in [0, 1]$ and then choose $x = P^{-1}(u)$. For any a < b, the probability that x is between a and b is equal to the probability that u is between P(a) and P(b) which equals P(b) - P(a) as desired. For the 2-dimensional Gaussian, one can generate a point in polar coordinates by choosing angle θ uniform in $[0, 2\pi]$ and radius $r = \sqrt{-2\ln(u)}$ where u is uniform random in [0, 1]. This is called the Box-Muller transform.

mean and variance σ^2 in each coordinate has density function

$$p(\mathbf{x}) = \frac{1}{\left(2\pi\right)^{d/2} \sigma^d} \exp\left(-\frac{|\mathbf{x}|^2}{2\sigma^2}\right).$$

The value of the density is maximum at the origin, but there is very little volume there. When $\sigma^2 = 1$, integrating the probability density over a unit ball centered at the origin yields almost zero mass since the volume of such a ball is negligible. In fact, one needs to increase the radius of the ball to nearly \sqrt{d} before there is a significant volume and hence significant probability mass. If one increases the radius much beyond \sqrt{d} , the integral barely increases even though the volume increases since the probability density is dropping off at a much higher rate. The following theorem formally states that nearly all the probability is concentrated in a thin annulus of width O(1) at radius \sqrt{d} .

Theorem 2.9 (Gaussian Annulus Theorem) For a d-dimensional spherical Gaussian with unit variance in each direction, for any $\beta \leq \sqrt{d}$, all but at most $3e^{-c\beta^2}$ of the probability mass lies within the annulus $\sqrt{d} - \beta \leq |\mathbf{x}| \leq \sqrt{d} + \beta$, where c is a fixed positive constant.

For a high-level intuition, note that $E(|\mathbf{x}|^2) = \sum_{i=1}^d E(x_i^2) = dE(x_1^2) = d$, so the mean squared distance of a point from the center is d. The Gaussian Annulus Theorem says that the points are tightly concentrated. We call the square root of the mean squared distance, namely \sqrt{d} , the radius of the Gaussian.

To prove the Gaussian Annulus Theorem we make use of a tail inequality for sums of independent random variables of bounded moments (Theorem 12.5).

Proof (Gaussian Annulus Theorem): Let $\mathbf{y} = (y_1, y_2, \dots, y_d)$ be a point selected from a unit variance Gaussian centered at the origin, and let $r = |\mathbf{y}|$. If $|r - \sqrt{d}| \ge \beta$, then multiplying both sides by $r + \sqrt{d}$ gives $|r^2 - d| \ge \beta(r + \sqrt{d}) \ge \beta\sqrt{d}$. So, it suffices to bound the probability that $|r^2 - d| \ge \beta\sqrt{d}$.

Rewrite $r^2 - d = (y_1^2 + \ldots + y_d^2) - d = (y_1^2 - 1) + \ldots + (y_d^2 - 1)$ and perform a change of variables: $x_i = y_i^2 - 1$. We want to bound the probability that $|x_1 + \ldots + x_d| \ge \beta \sqrt{d}$. Notice that $E(x_i) = E(y_i^2) - 1 = 0$. To apply Theorem 12.5, we need to bound the s^{th} moments of x_i .

For $|y_i| \leq 1$, $|x_i|^s \leq 1$ and for $|y_i| \geq 1$, $|x_i|^s \leq |y_i|^{2s}$. Thus

$$|E(x_i^s)| = E(|x_i|^s) \le E(1+y_i^{2s}) = 1 + E(y_i^{2s})$$
$$= 1 + \sqrt{\frac{2}{\pi}} \int_0^\infty y^{2s} e^{-y^2/2} dy$$

Using the substitution $2z = y^2$,

$$\begin{split} |E(x_i^s)| &= 1 + \frac{1}{\sqrt{\pi}} \int_0^\infty 2^s z^{s - (1/2)} e^{-z} dz \\ &\leq 2^s s!. \end{split}$$

The last inequality is from the Gamma integral.

Since $E(x_i) = 0$, $Var(x_i) = E(x_i^2) \le 2^2 2 = 8$. Unfortunately, we do not have $|E(x_i^s)| \le 8s!$ as required in Theorem 12.5. To fix this problem, perform one more change of variables, using $w_i = x_i/2$. Then, $Var(w_i) \le 2$ and $|E(w_i^s)| \le 2s!$, and our goal is now to bound the probability that $|w_1 + \ldots + w_d| \ge \frac{\beta\sqrt{d}}{2}$. Applying Theorem 12.5 where $\sigma^2 = 2$ and n = d, this occurs with probability less than or equal to $3e^{-\frac{\beta^2}{96}}$.

In the next sections we will see several uses of the Gaussian Annulus Theorem.

2.7 Random Projection and Johnson-Lindenstrauss Lemma

One of the most frequently used subroutines in tasks involving high dimensional data is nearest neighbor search. In nearest neighbor search we are given a database of n points in \mathbf{R}^d where n and d are usually large. The database can be preprocessed and stored in an efficient data structure. Thereafter, we are presented "query" points in \mathbf{R}^d and are to find the nearest or approximately nearest database point to the query point. Since the number of queries is often large, query time (time to answer a single query) should be very small (ideally a small function of log n and log d), whereas preprocessing time could be larger (a polynomial function of n and d). For this and other problems, dimension reduction, where one projects the database points to a k dimensional space with $k \ll d$ (usually dependent on log d) can be very useful so long as the relative distances between points are approximately preserved. We will see using the Gaussian Annulus Theorem that such a projection indeed exists and is simple.

The projection $f : \mathbf{R}^d \to \mathbf{R}^k$ that we will examine (in fact, many related projections are known to work as well) is the following. Pick k vectors $\mathbf{u_1}, \mathbf{u_2}, \ldots, \mathbf{u_k}$, independently from the Gaussian distribution $\frac{1}{(2\pi)^{d/2}} \exp(-|\mathbf{x}|^2/2)$. For any vector \mathbf{v} , define the projection $f(\mathbf{v})$ by:

$$f(\mathbf{v}) = (\mathbf{u_1} \cdot \mathbf{v}, \mathbf{u_2} \cdot \mathbf{v}, \dots, \mathbf{u_k} \cdot \mathbf{v}).$$

The projection $f(\mathbf{v})$ is the vector of dot products of \mathbf{v} with the \mathbf{u}_i . We will show that with high probability, $|f(\mathbf{v})| \approx \sqrt{k}|\mathbf{v}|$. For any two vectors $\mathbf{v_1}$ and $\mathbf{v_2}$, $f(\mathbf{v_1} - \mathbf{v_2}) = f(\mathbf{v_1}) - f(\mathbf{v_2})$. Thus, to estimate the distance $|\mathbf{v_1} - \mathbf{v_2}|$ between two vectors $\mathbf{v_1}$ and $\mathbf{v_2}$ in \mathbf{R}^d , it suffices to compute $|f(\mathbf{v_1}) - f(\mathbf{v_2})| = |f(\mathbf{v_1} - \mathbf{v_2})|$ in the k dimensional space since the factor of \sqrt{k} is known and one can divide by it. The reason distances increase when we project to a lower dimensional space is that the vectors $\mathbf{u_i}$ are not unit length. Also notice that the vectors $\mathbf{u_i}$ are not orthogonal. If we had required them to be orthogonal, we would have lost statistical independence. **Theorem 2.10 (The Random Projection Theorem)** Let \mathbf{v} be a fixed vector in \mathbf{R}^d and let f be defined as above. Then there exists constant c > 0 such that for $\varepsilon \in (0, 1)$,

$$Prob\left(\left|\left|f(\mathbf{v})\right| - \sqrt{k}|\mathbf{v}|\right| \ge \varepsilon\sqrt{k}|\mathbf{v}|\right) \le 3e^{-ck\varepsilon^2},$$

where the probability is taken over the random draws of vectors \mathbf{u}_{i} used to construct f.

Proof: By scaling both sides by $|\mathbf{v}|$, we may assume that $|\mathbf{v}| = 1$. The sum of independent normally distributed real variables is also normally distributed where the mean and variance are the sums of the individual means and variances. Since $\mathbf{u}_i \cdot \mathbf{v} = \sum_{j=1}^d u_{ij}v_j$, the random variable $\mathbf{u}_i \cdot \mathbf{v}$ has Gaussian density with zero mean and variance equal to $\sum_{j=1}^d v_j^2 = |\mathbf{v}|^2 = 1$. (Since u_{ij} has variance one and the v_j is a constant, the variance of $u_{ij}v_j$ is v_j^2 .) Since $\mathbf{u}_1 \cdot \mathbf{v}, \mathbf{u}_2 \cdot \mathbf{v}, \dots, \mathbf{u}_k \cdot \mathbf{v}$ are independent, the theorem follows from the Gaussian Annulus Theorem (Theorem 2.9) with k = d.

The random projection theorem establishes that the probability of the length of the projection of a single vector differing significantly from its expected value is exponentially small in k, the dimension of the target subspace. By a union bound, the probability that any of $O(n^2)$ pairwise differences $|\mathbf{v_i} - \mathbf{v_j}|$ among n vectors $\mathbf{v_1}, \ldots, \mathbf{v_n}$ differs significantly from their expected values is small, provided $k \geq \frac{3}{c\varepsilon^2} \ln n$. Thus, this random projection preserves all relative pairwise distances between points in a set of n points with high probability. This is the content of the Johnson-Lindenstrauss Lemma.

Theorem 2.11 (Johnson-Lindenstrauss Lemma) For any $0 < \varepsilon < 1$ and any integer n, let $k \geq \frac{3}{c\varepsilon^2} \ln n$ for c as in Theorem 2.9. For any set of n points in \mathbb{R}^d , the random projection $f: \mathbb{R}^d \to \mathbb{R}^k$ defined above has the property that for all pairs of points $\mathbf{v_i}$ and $\mathbf{v_j}$, with probability at least 1 - 1.5/n,

$$(1-\varepsilon)\sqrt{k} |\mathbf{v}_{\mathbf{i}} - \mathbf{v}_{\mathbf{j}}| \le |f(\mathbf{v}_{\mathbf{i}}) - f(\mathbf{v}_{\mathbf{j}})| \le (1+\varepsilon)\sqrt{k} |\mathbf{v}_{\mathbf{i}} - \mathbf{v}_{\mathbf{j}}|.$$

Proof: Applying the Random Projection Theorem (Theorem 2.10), for any fixed $\mathbf{v_i}$ and $\mathbf{v_j}$, the probability that $|f(\mathbf{v_i} - \mathbf{v_j})|$ is outside the range

$$\left[(1-\varepsilon)\sqrt{k}|\mathbf{v_i}-\mathbf{v_j}|,(1+\varepsilon)\sqrt{k}|\mathbf{v_i}-\mathbf{v_j}|\right]$$

is at most $3e^{-ck\varepsilon^2} \leq 3/n^3$ for $k \geq \frac{3\ln n}{c\varepsilon^2}$. Since there are $\binom{n}{2} < n^2/2$ pairs of points, by the union bound, the probability that any pair has a large distortion is less than $\frac{3}{2n}$.

Remark: It is important to note that the conclusion of Theorem 2.11 asserts for all $\mathbf{v_i}$ and $\mathbf{v_j}$, not just for most of them. The weaker assertion for most $\mathbf{v_i}$ and $\mathbf{v_j}$ is typically less useful, since our algorithm for a problem such as nearest-neighbor search might return one of the bad pairs of points. A remarkable aspect of the theorem is that the number of dimensions in the projection is only dependent logarithmically on n. Since k is often much less than d, this is called a dimension reduction technique. In applications, the dominant term is typically the inverse square dependence on ε .

For the nearest neighbor problem, if the database has n_1 points and n_2 queries are expected during the lifetime of the algorithm, take $n = n_1 + n_2$ and project the database to a random k-dimensional space, for k as in Theorem 2.11. On receiving a query, project the query to the same subspace and compute nearby database points. The Johnson Lindenstrauss Theorem says that with high probability this will yield the right answer whatever the query. Note that the exponentially small in k probability was useful here in making k only dependent on $\ln n$, rather than n.

2.8 Separating Gaussians

Mixtures of Gaussians are often used to model heterogeneous data coming from multiple sources. For example, suppose we are recording the heights of individuals age 20-30 in a city. We know that on average, men tend to be taller than women, so a natural model would be a Gaussian mixture model $p(x) = w_1p_1(x) + w_2p_2(x)$, where $p_1(x)$ is a Gaussian density representing the typical heights of women, $p_2(x)$ is a Gaussian density representing the typical heights of men, and w_1 and w_2 are the mixture weights representing the proportion of women and men in the city. The parameter estimation problem for a mixture model is the problem: given access to samples from the overall density p (e.g., heights of people in the city, but without being told whether the person with that height is male or female), reconstruct the parameters for the distribution (e.g., good approximations to the means and variances of p_1 and p_2 , as well as the mixture weights).

There are taller women and shorter men, so even if one solved the parameter estimation problem for heights perfectly, given a data point (a height) one couldn't necessarily tell which population it came from (male or female). In this section, we will look at a problem that is in some ways easier and some ways harder than this problem of heights. It will be harder in that we will be interested in a mixture of two Gaussians in highdimensions (as opposed to the d = 1 case of heights). But it will be easier in that we will assume the means are quite well-separated compared to the variances. Specifically, our focus will be on a mixture of two spherical unit-variance Gaussians whose means are separated by a distance $\Omega(d^{1/4})$. We will show that at this level of separation, we can with high probability uniquely determine which Gaussian each data point came from. The algorithm to do so will actually be quite simple. Calculate the distance between all pairs of points. Points whose distance apart is smaller are from the same Gaussian, whereas points whose distance is larger are from different Gaussians. Later, we will see that with more sophisticated algorithms, even a separation of $\Omega(1)$ suffices.

First, consider just one spherical unit-variance Gaussian centered at the origin. From Theorem 2.9, most of its probability mass lies on an annulus of width O(1) at radius \sqrt{d} . Also $e^{-|\mathbf{x}|^2/2} = \prod_i e^{-x_i^2/2}$ and almost all of the mass is within the slab { $\mathbf{x} \mid -c \leq x_1 \leq c$ }, for $c \in O(1)$. Pick a point \mathbf{x} from this Gaussian. After picking \mathbf{x} , rotate the coordinate system to make the first axis align with \mathbf{x} . Independently pick a second point \mathbf{y} from this Gaussian. The fact that almost all of the probability mass of the Gaussian is within

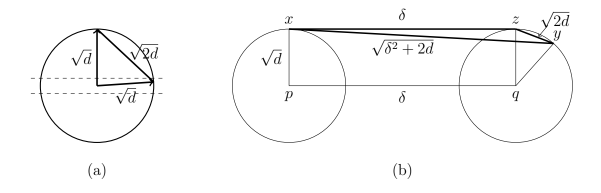


Figure 2.4: (a) indicates that two randomly chosen points in high dimension are surely almost nearly orthogonal. (b) indicates that the distance between a pair of random points from two different unit balls approximating the annuli of two Gaussians.

the slab $\{\mathbf{x} \mid -c \leq x_1 \leq c, c \in O(1)\}$ at the equator implies that **y**'s component along **x**'s direction is O(1) with high probability. Thus, **y** is nearly perpendicular to **x**. So, $|\mathbf{x} - \mathbf{y}| \approx \sqrt{|\mathbf{x}|^2 + |\mathbf{y}|^2}$. See Figure 2.4(a). More precisely, since the coordinate system has been rotated so that **x** is at the North Pole, $\mathbf{x} = (\sqrt{d} \pm O(1), 0, \dots, 0)$. Since **y** is almost on the equator, further rotate the coordinate system so that the component of **y** that is perpendicular to the axis of the North Pole is in the second coordinate. Then $\mathbf{y} = (O(1), \sqrt{d} \pm O(1), 0, \dots, 0)$. Thus,

$$(\mathbf{x} - \mathbf{y})^2 = d \pm O(\sqrt{d}) + d \pm O(\sqrt{d}) = 2d \pm O(\sqrt{d})$$

and $|\mathbf{x} - \mathbf{y}| = \sqrt{2d} \pm O(1)$ with high probability.

Consider two spherical unit variance Gaussians with centers \mathbf{p} and \mathbf{q} separated by a distance Δ . The distance between a randomly chosen point \mathbf{x} from the first Gaussian and a randomly chosen point \mathbf{y} from the second is close to $\sqrt{\Delta^2 + 2d}$, since $\mathbf{x} - \mathbf{p}, \mathbf{p} - \mathbf{q}$, and $\mathbf{q} - \mathbf{y}$ are nearly mutually perpendicular. Pick \mathbf{x} and rotate the coordinate system so that \mathbf{x} is at the North Pole. Let \mathbf{z} be the North Pole of the ball approximating the second Gaussian. Now pick \mathbf{y} . Most of the mass of the second Gaussian is within O(1) of the equator perpendicular to $\mathbf{q} - \mathbf{z}$. Also, most of the mass of each Gaussian is within distance O(1) of the respective equators perpendicular to the line $\mathbf{q} - \mathbf{p}$. See Figure 2.4 (b). Thus,

$$|\mathbf{x} - \mathbf{y}|^2 \approx \Delta^2 + |\mathbf{z} - \mathbf{q}|^2 + |\mathbf{q} - \mathbf{y}|^2$$
$$= \Delta^2 + 2d \pm O(\sqrt{d})).$$

To ensure that the distance between two points picked from the same Gaussian are closer to each other than two points picked from different Gaussians requires that the upper limit of the distance between a pair of points from the same Gaussian is at most the lower limit of distance between points from different Gaussians. This requires that $\sqrt{2d} + O(1) \leq \sqrt{2d + \Delta^2} - O(1)$ or $2d + O(\sqrt{d}) \leq 2d + \Delta^2$, which holds when $\Delta \in \omega(d^{1/4})$. Thus, mixtures of spherical Gaussians can be separated in this way, provided their centers are separated by $\omega(d^{1/4})$. If we have *n* points and want to correctly separate all of them with high probability, we need our individual high-probability statements to hold with probability 1 - 1/poly(n),² which means our O(1) terms from Theorem 2.9 become $O(\sqrt{\log n})$. So we need to include an extra $O(\sqrt{\log n})$ term in the separation distance.

Algorithm for separating points from two Gaussians: Calculate all pairwise distances between points. The cluster of smallest pairwise distances must come from a single Gaussian. Remove these points. The remaining points come from the second Gaussian.

One can actually separate Gaussians where the centers are much closer. In the next chapter we will use singular value decomposition to separate a mixture of two Gaussians when their centers are separated by a distance O(1).

2.9 Fitting a Single Spherical Gaussian to Data

Given a set of sample points, $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_n}$, in a *d*-dimensional space, we wish to find the spherical Gaussian that best fits the points. Let *F* be the unknown Gaussian with mean $\boldsymbol{\mu}$ and variance σ^2 in each direction. The probability density for picking these points when sampling according to *F* is given by

$$c \exp\left(-\frac{(\mathbf{x_1}-\boldsymbol{\mu})^2+(\mathbf{x_2}-\boldsymbol{\mu})^2+\cdots+(\mathbf{x_n}-\boldsymbol{\mu})^2}{2\sigma^2}\right)$$

where the normalizing constant c is the reciprocal of $\left[\int e^{-\frac{|\mathbf{x}-\boldsymbol{\mu}|^2}{2\sigma^2}}dx\right]^n$. In integrating from $-\infty$ to ∞ , one can shift the origin to $\boldsymbol{\mu}$ and thus c is $\left[\int e^{-\frac{|\mathbf{x}|^2}{2\sigma^2}}dx\right]^{-n} = \frac{1}{(2\pi)^{\frac{n}{2}}}$ and is independent of $\boldsymbol{\mu}$.

The Maximum Likelihood Estimator (MLE) of F, given the samples $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_n}$, is the F that maximizes the above probability density.

Lemma 2.12 Let $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n\}$ be a set of *n* d-dimensional points. Then $(\mathbf{x}_1 - \boldsymbol{\mu})^2 + (\mathbf{x}_2 - \boldsymbol{\mu})^2 + \dots + (\mathbf{x}_n - \boldsymbol{\mu})^2$ is minimized when $\boldsymbol{\mu}$ is the centroid of the points $\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n$, namely $\boldsymbol{\mu} = \frac{1}{n} (\mathbf{x}_1 + \mathbf{x}_2 + \dots + \mathbf{x}_n)$.

Proof: Setting the gradient of $(\mathbf{x}_1 - \boldsymbol{\mu})^2 + (\mathbf{x}_2 - \boldsymbol{\mu})^2 + \dots + (\mathbf{x}_n - \boldsymbol{\mu})^2$ with respect $\boldsymbol{\mu}$ to zero yields

$$-2\left(\mathbf{x}_{1}-\boldsymbol{\mu}\right)-2\left(\mathbf{x}_{2}-\boldsymbol{\mu}\right)-\cdots-2\left(\mathbf{x}_{n}-\boldsymbol{\mu}\right)=0.$$

Solving for $\boldsymbol{\mu}$ gives $\boldsymbol{\mu} = \frac{1}{n}(\mathbf{x}_1 + \mathbf{x}_2 + \cdots + \mathbf{x}_n).$

²poly(n) means bounded by a polynomial in n.

To determine the maximum likelihood estimate of σ^2 for F, set $\boldsymbol{\mu}$ to the true centroid. Next, show that σ is set to the standard deviation of the sample. Substitute $\nu = \frac{1}{2\sigma^2}$ and $a = (\mathbf{x}_1 - \boldsymbol{\mu})^2 + (\mathbf{x}_2 - \boldsymbol{\mu})^2 + \dots + (\mathbf{x}_n - \boldsymbol{\mu})^2$ into the formula for the probability of picking the points $\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_n$. This gives

$$\frac{e^{-a\nu}}{\left[\int\limits_{x}e^{-x^{2}\nu}dx\right]^{n}}$$

Now, a is fixed and ν is to be determined. Taking logs, the expression to maximize is

$$-a\nu - n\ln\left[\int\limits_{x}e^{-\nu x^{2}}dx
ight].$$

To find the maximum, differentiate with respect to ν , set the derivative to zero, and solve for σ . The derivative is

$$-a + n \frac{\int\limits_{x} |x|^2 e^{-\nu x^2} dx}{\int\limits_{x} e^{-\nu x^2} dx}.$$

Setting $y = |\sqrt{\nu} \mathbf{x}|$ in the derivative, yields

$$-a+rac{n}{
u}rac{\int\limits_{y}y^{2}e^{-y^{2}}dy}{\int\limits_{y}e^{-y^{2}}dy}.$$

Since the ratio of the two integrals is the expected distance squared of a *d*-dimensional spherical Gaussian of standard deviation $\frac{1}{\sqrt{2}}$ to its center, and this is known to be $\frac{d}{2}$, we get $-a + \frac{nd}{2\nu}$. Substituting σ^2 for $\frac{1}{2\nu}$ gives $-a + nd\sigma^2$. Setting $-a + nd\sigma^2 = 0$ shows that the maximum occurs when $\sigma = \frac{\sqrt{a}}{\sqrt{nd}}$. Note that this quantity is the square root of the average coordinate distance squared of the samples to their mean, which is the standard deviation of the sample. Thus, we get the following lemma.

Lemma 2.13 The maximum likelihood spherical Gaussian for a set of samples is the Gaussian with center equal to the sample mean and standard deviation equal to the standard deviation of the sample from the true mean.

Let $\mathbf{x}_1, \mathbf{x}_2, \ldots, \mathbf{x}_n$ be a sample of points generated by a Gaussian probability distribution. Then $\boldsymbol{\mu} = \frac{1}{n}(\mathbf{x}_1 + \mathbf{x}_2 + \cdots + \mathbf{x}_n)$ is an unbiased estimator of the expected value of the distribution. However, if in estimating the variance from the sample set, we use the estimate of the expected value rather than the true expected value, we will not get an unbiased estimate of the variance, since the sample mean is not independent of the sample set. One should use $\tilde{\boldsymbol{\mu}} = \frac{1}{n-1}(\mathbf{x}_1 + \mathbf{x}_2 + \cdots + \mathbf{x}_n)$ when estimating the variance. See Section ?? of the appendix.

2.10 Bibliographic Notes

The word vector model was introduced by Salton [SWY75]. There is vast literature on the Gaussian distribution, its properties, drawing samples according to it, etc. The reader can choose the level and depth according to his/her background. The Master Tail Bounds theorem and the derivation of Chernoff and other inequalities from it are from [Kan09]. The original proof of the Random Projection Theorem by Johnson and Lindenstrauss was complicated. Several authors used Gaussians to simplify the proof. The proof here is due to Dasgupta and Gupta [DG99]. See [Vem04] for details and applications of the theorem. [MU05] and [MR95b] are text books covering a lot of the material touched upon here.

2.11 Exercises

Exercise 2.1

- 1. Let x and y be independent random variables with uniform distribution in [0,1]. What is the expected value E(x), $E(x^2)$, E(x-y), E(xy), and $E(x-y)^2$?
- 2. Let x and y be independent random variables with uniform distribution in $\left[-\frac{1}{2}, \frac{1}{2}\right]$. What is the expected value E(x), $E(x^2)$, E(x-y), E(xy), and $E(x-y)^2$?
- 3. What is the expected squared distance between two points generated at random inside a unit d-dimensional cube?

Exercise 2.2 Randomly generate 30 points inside the cube $\left[-\frac{1}{2}, \frac{1}{2}\right]^{100}$ and plot distance between points and the angle between the vectors from the origin to the points for all pairs of points.

Exercise 2.3 Show that Markov's inequality is tight by showing the following:

- 1. For each a = 2, 3, and 4 give a probability distribution p(x) for a nonnegative random variable x where $Prob(x \ge a) = \frac{E(x)}{a}$.
- 2. For arbitrary $a \ge 1$ give a probability distribution for a nonnegative random variable x where $\operatorname{Prob}(x \ge a) = \frac{E(x)}{a}$.

Exercise 2.4 Give a probability distribution p(x) and a value b for which Chebyshev's inequality is tight and a probability distribution and value of b for which it is not tight.

Exercise 2.5 Consider the probability density function p(x) = 0 for x < 1 and $p(x) = c\frac{1}{x^4}$ for $x \ge 1$.

- 1. What should c be to make p a legal probability density function?
- 2. Generate 100 random samples from this distribution. How close is the average of the samples to the expected value of x?

Exercise 2.6 Let G be a d-dimensional spherical Gaussian with variance $\frac{1}{2}$ centered at the origin. Derive the expected squared distance to the origin.

Exercise 2.7 Consider drawing a random point \mathbf{x} on the surface of the unit sphere in \mathbb{R}^d . What is the variance of x_1 (the first coordinate of \mathbf{x})? See if you can give an argument without doing any integrals.

Exercise 2.8 How large must ε be for 99% of the volume of a d-dimensional unit-radius ball to lie in the shell of ε -thickness at the surface of the ball?

Exercise 2.9 A 3-dimensional cube has vertices, edges, and faces. In a d-dimensional cube, these components are called faces. A vertex is a 0-dimensional face, an edge a 1-dimensional face, etc.

- 1. For $0 \le k \le d$, how many k-dimensional faces does a d-dimensional cube have?
- 2. What is the total number of faces of all dimensions? The d-dimensional face is the cube itself which you can include in your count.
- 3. What is the surface area of a unit cube in d-dimensions (a unit cube has side-length 1 in each dimension)?
- 4. What is the surface area of the cube if the length of each side was 2?
- 5. Prove that the volume of a unit cube is close to its surface.

Exercise 2.10 Consider the portion of the surface area of a unit radius, 3-dimensional ball with center at the origin that lies within a circular cone whose vertex is at the origin. What is the formula for the incremental unit of area when using polar coordinates to integrate the portion of the surface area of the ball that is lying inside the circular cone? What is the formula for the integral? What is the value of the integral if the angle of the cone is 36° ? The angle of the cone is measured from the axis of the cone to a ray on the surface of the cone.

Exercise 2.11 For what value of d does the volume, V(d), of a d-dimensional unit ball take on its maximum? Hint: Consider the ratio V(d)

Hint: Consider the ratio $\frac{V(d)}{V(d-1)}$.

Exercise 2.12 How does the volume of a ball of radius two behave as the dimension of the space increases? What if the radius was larger than two but a constant independent of d? What function of d would the radius need to be for a ball of radius r to have approximately constant volume as the dimension increases?

Exercise 2.13 If $\lim_{d\to\infty} V(d) = 0$, the volume of a d-dimensional ball for sufficiently large d must be less than V(3). How can this be if the d-dimensional ball contains the three dimensional ball?

Exercise 2.14 Consider a unit radius, circular cylinder in 3-dimensions of height one. The top of the cylinder could be an horizontal plane or half of a circular ball. Consider these two possibilities for a unit radius, circular cylinder in 4-dimensions. In 4-dimensions the horizontal plane is 3-dimensional and the half circular ball is 4-dimensional. In each of the two cases, what is the surface area of the top face of the cylinder? You can use V(d) for the volume of a unit radius, d-dimension ball and A(d) for the surface area of a unit radius, d-dimension ball and A(d) for the surface area of the top face of the surface area of the surface area of a unit radius, d-dimensional ball. An infinite length, unit radius, circular cylinder in 4-dimensions would be the set $\{(x_1, x_2, x_3, x_4) | x_2^2 + x_3^2 + x_4^2 \leq 1\}$ where the coordinate x_1 is the axis. Exercise 2.15 Given a d-dimensional circular cylinder of radius r and height h

- 1. What is the surface area in terms of V(d) and A(d)?
- 2. What is the volume?

Exercise 2.16 Write a recurrence relation for V(d) in terms of V(d-1) by integrating using an incremental unit that is a disk of thickness dr.

Exercise 2.17 Verify the formula $V(d) = 2 \int_0^1 V(d-1)(1-x_1^2)^{\frac{d-1}{2}} dx_1$ for d=2 and d=3 by integrating and comparing with $V(2) = \pi$ and $V(3) = \frac{4}{3}\pi$

Exercise 2.18 Consider a unit ball A centered at the origin and a unit ball B whose center is at distance s from the origin. Suppose that a random point x is drawn from the mixture distribution: "with probability 1/2, draw at random from A; with probability 1/2, draw at random from B". Show that a separation $s \gg 1/\sqrt{d-1}$ is sufficient so that $Prob(x \in A \cap B) = o(1)$; i.e., for any $\epsilon > 0$ there exists c such that if $s \ge c/\sqrt{d-1}$ then $Prob(x \in A \cap B) < \epsilon$. In other words, this extent of separation means that nearly all of the mixture distribution is identifiable.

Exercise 2.19 Prove that $1 + x \le e^x$ for all real x. For what values of x is the approximation $1 + x \approx e^x$ within 0.01?

Exercise 2.20 Consider the upper hemisphere of a unit-radius ball in d-dimensions. What is the height of the maximum volume cylinder that can be placed entirely inside the hemisphere? As you increase the height of the cylinder, you need to reduce the cylinder's radius so that it will lie entirely within the hemisphere.

Exercise 2.21 What is the volume of the maximum size d-dimensional hypercube that can be placed entirely inside a unit radius d-dimensional ball?

Exercise 2.22 For a 1,000-dimensional unit-radius ball centered at the origin, what fraction of the volume of the upper hemisphere is above the plane $x_1 = 0.1$? Above the plane $x_1 = 0.01$?

Exercise 2.23 Calculate the ratio of area above the plane $x_1 = \epsilon$ to the area of the upper hemisphere of a unit radius ball in d-dimensions for $\epsilon = 0.01, 0.02, 0.03, 0.04, 0.05$ and for d = 100 and d = 1,000. Also calculate the ratio for $\epsilon = 0.001$ and d = 1,000.

Exercise 2.24 Let $\{\mathbf{x} \mid |\mathbf{x}| \leq 1\}$ be a d-dimensional, unit radius ball centered at the origin. What fraction of the volume is the set $\{(x_1, x_2, \ldots, x_d) \mid |x_i| \leq \frac{1}{\sqrt{d}}\}$?

Exercise 2.25 Almost all of the volume of a ball in high dimensions lies in a narrow slice of the ball at the equator. However, the narrow slice is determined by the point on the surface of the ball that is designated the North Pole. Explain how this can be true if several different locations are selected for the location of the North Pole giving rise to different equators.

Exercise 2.26 Explain how the volume of a ball in high dimensions can simultaneously be in a narrow slice at the equator and also be concentrated in a narrow annulus at the surface of the ball.

Exercise 2.27 Generate 500 points uniformly at random on the surface of a unit-radius ball in 50 dimensions. Then randomly generate five additional points. For each of the five new points, calculate a narrow band of width $\frac{2}{\sqrt{50}}$ at the equator, assuming the point was the North Pole. How many of the 500 points are in each band corresponding to one of the five equators? How many of the points are in all five bands? How wide do the bands need to be for all points to be in all five bands?

Exercise 2.28 Consider a slice of a 100-dimensional ball that lies between two parallel planes, each equidistant from the equator and perpendicular to the line from the North Pole to the South Pole. What percentage of the distance from the center of the ball to the poles must the planes be to contain 95% of the surface area?

Exercise 2.29 Place 100 points at random on a d-dimensional unit-radius ball. Assume d is large. Pick a random vector and let it define two parallel hyperplanes on opposite sides of the origin that are equal distance from the origin. How far apart can the hyperplanes be moved and still have the probability that none of the n points lands between them be at least .99?

Exercise 2.30 Consider two random vectors in a high-dimensional space. Assume the vectors have been normalized so that their lengths are one and thus the points lie on a unit ball. Assume one of the vectors is the North pole. Prove that the ratio of the surface area of a cone, with axis at the North Pole of fixed angle say 45° to the area of a hemisphere, goes to zero as the dimension increases. Thus, the probability that the angle between two random vectors is at most 45° goes to zero. How does this relate to the result that most of the volume is near the equator?

Exercise 2.31 Project the volume of a d-dimensional ball of radius \sqrt{d} onto a line through the center. For large d, give an intuitive argument that the projected volume should behave like a Gaussian.

Exercise 2.32

- 1. Write a computer program that generates n points uniformly distributed over the surface of a unit-radius d-dimensional ball.
- 2. Generate 200 points on the surface of a sphere in 50 dimensions.
- 3. Create several random lines through the origin and project the points onto each line. Plot the distribution of points on each line.
- 4. What does your result from (3) say about the surface area of the sphere in relation to the lines, i.e., where is the surface area concentrated relative to each line?

Exercise 2.33 If one generates points in d-dimensions with each coordinate a unit variance Gaussian, the points will approximately lie on the surface of a sphere of radius \sqrt{d} .

- 1. What is the distribution when the points are projected onto a random line through the origin?
- 2. If one uses a Gaussian with variance four, where in d-space will the points lie?

Exercise 2.34 Randomly generate a 100 points on the surface of a sphere in 3-dimensions and in 100-dimensions. Create a histogram of all distances between the pairs of points in both cases.

Exercise 2.35 We have claimed that a randomly generated point on a ball lies near the equator of the ball, independent of the point picked to be the North Pole. Is the same claim true for a randomly generated point on a cube? To test this claim, randomly generate ten ± 1 valued vectors in 128 dimensions. Think of these ten vectors as ten choices for the North Pole. Then generate some additional ± 1 valued vectors. To how many of the original vectors is each of the new vectors close to being perpendicular; that is, how many of the equators is each new vector close to?

Exercise 2.36 Project the vertices of a high-dimensional cube onto a line from (0, 0, ..., 0) to (1, 1, ..., 1). Argue that the "density" of the number of projected points (per unit distance) varies roughly as a Gaussian with variance O(1) with the mid-point of the line as center.

Exercise 2.37 Define the equator of a d-dimensional unit cube to be the hyperplane $\left\{ \mathbf{x} \middle| \sum_{i=1}^{d} x_i = \frac{d}{2} \right\}$.

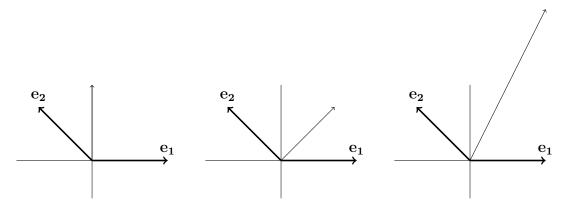
- 1. Are the vertices of a unit cube concentrated close to the equator?
- 2. Is the volume of a unit cube concentrated close to the equator?
- 3. Is the surface area of a unit cube concentrated close to the equator?

Exercise 2.38 Let x be a random variable with probability density $\frac{1}{4}$ for $0 \le x \le 4$ and zero elsewhere.

- 1. Use Markov's inequality to bound the probability that x > 3.
- 2. Make use of $Prob(|x| > a) = Prob(x^2 > a^2)$ to get a tighter bound.
- 3. What is the bound using $Prob(|x| > a) = Prob(x^r > a^r)$?

Exercise 2.39 Consider the probability distribution $p(x = 0) = 1 - \frac{1}{a}$ and $p(x = a) = \frac{1}{a}$. Plot the probability that x is greater than or equal to a as a function of a for the bound given by Markov's inequality and by Markov's inequality applied to x^2 and x^4 . **Exercise 2.40** Consider a non orthogonal basis $\mathbf{e_1}, \mathbf{e_2}, \ldots, \mathbf{e_d}$. The $\mathbf{e_i}$ are a set of linearly independent unit vectors that span the space.

- 1. Prove that the representation of any vector in this basis is unique.
- 2. Calculate the squared length of $\mathbf{z} = (\frac{\sqrt{2}}{2}, 1)_e$ where \mathbf{z} is expressed in the basis $\mathbf{e_1} = (1,0)$ and $\mathbf{e_2} = (-\frac{\sqrt{2}}{2}, \frac{\sqrt{2}}{2})$
- 3. If $\mathbf{y} = \sum_{i} a_i \mathbf{e}_i$ and $\mathbf{z} = \sum_{i} b_i \mathbf{e}_i$, with $0 < a_i < b_i$, is it necessarily true that the length of \mathbf{z} is greater than the length of \mathbf{y} ? Why or why not?
- 4. Consider the basis $\mathbf{e_1} = (1,0)$ and $\mathbf{e_2} = (-\frac{\sqrt{2}}{2},\frac{\sqrt{2}}{2})$.
 - (a) What is the representation of the vector (0,1) in the basis $(\mathbf{e_1},\mathbf{e_2})$.
 - (b) What is the representation of the vector $(\frac{\sqrt{2}}{2}, \frac{\sqrt{2}}{2})$?
 - (c) What is the representation of the vector (1,2)?



Exercise 2.41 Generate 20 points uniformly at random on a 900-dimensional sphere of radius 30. Calculate the distance between each pair of points. Then, select a method of projection and project the data onto subspaces of dimension k=100, 50, 10, 5, 4, 3, 2, 1 and calculate the difference between \sqrt{k} times the original distances and the new pair-wise distances. For each value of k what is the maximum difference as a percent of \sqrt{k} .

Exercise 2.42 In d-dimensions there are exactly d-unit vectors that are pairwise orthogonal. However, if you wanted a set of vectors that were almost orthogonal you might squeeze in a few more. For example, in 2-dimensions if almost orthogonal meant at least 45 degrees apart you could fit in three almost orthogonal vectors. Suppose you wanted to find 900 almost orthogonal vectors in 100 dimensions where almost orthogonal meant an angle of between 85 and 95 degrees. How would you generate such a set?

Hint: Consider projecting a 1,000 orthonormal 1,000-dimensional vectors to a random 100-dimensional space.

Exercise 2.43 Exercise 2.42 finds almost orthogonal vectors using the Johnson Lindenstrauss Theorem. One could also create almost orthogonal vectors by generating random Gaussian vectors. Compare the two results to see which does a better job.

Exercise 2.44 To preserve pairwise distances between n data points in d space, we projected to a random $O(\ln n/\varepsilon^2)$ dimensional space. To save time in carrying out the projection, we may try to project to a space spanned by sparse vectors, vectors with only a few nonzero entries. That is, choose say $O(\ln n/\varepsilon^2)$ vectors at random, each with 100 nonzero components and project to the space spanned by them. Will this work to preserve approximately all pairwise distances? Why?

Exercise 2.45 Suppose there is an object moving at constant velocity along a straight line. You receive the gps coordinates corrupted by Gaussian noise every minute. How do you estimate the current position?

Exercise 2.46

- 1. What is the maximum size rectangle that can be fitted under a unit variance Gaussian?
- 2. What rectangle best approximates a unit variance Gaussian if one measure goodness of fit by the symmetric difference of the Gaussian and the rectangle.

Exercise 2.47 Let x_1, x_2, \ldots, x_n be independent samples of a random variable **x** with mean μ and variance σ^2 . Let $m_s = \frac{1}{n} \sum_{i=1}^n x_i$ be the sample mean. Suppose one estimates the variance using the sample mean rather than the true mean, that is,

$$\sigma_s^2 = \frac{1}{n} \sum_{i=1}^n (x_i - m_s)^2$$

Prove that $E(\sigma_s^2) = \frac{n-1}{n}\sigma^2$ and thus one should have divided by n-1 rather than n.

Hint: First calculate the variance of the sample mean and show that $var(m_s) = \frac{1}{n}var(\mathbf{x})$. *Then calculate* $E(\sigma_s^2) = E[\frac{1}{n}\sum_{i=1}^n (x_i - m_s)^2]$ by replacing $x_i - m_s$ with $(x_i - m) - (m_s - m)$.

Exercise 2.48 Generate ten values by a Gaussian probability distribution with zero mean and variance one. What is the center determined by averaging the points? What is the variance? In estimating the variance, use both the real center and the estimated center. When using the estimated center to estimate the variance, use both n = 10 and n = 9. How do the three estimates compare?

Exercise 2.49 Suppose you want to estimate the unknown center of a Gaussian in dspace which has variance one in each direction. Show that $O(\log d/\varepsilon^2)$ random samples from the Gaussian are sufficient to get an estimate **m** of the true center μ , so that with probability at least 99/100,

$$|\mu - \mathbf{m}|_{\infty} \leq \varepsilon.$$

How many samples are sufficient to ensure that

$$|\mu - \mathbf{m}| \le \varepsilon$$
?

Exercise 2.50 Use the probability distribution $\frac{1}{3\sqrt{2\pi}}e^{-\frac{1}{2}\frac{(x-5)^2}{9}}$ to generate ten points.

- (a) From the ten points estimate μ . How close is the estimate of μ to the true mean of 5?
- (b) Using the true mean of 5, estimate σ^2 by the formula $\sigma^2 = \frac{1}{10} \sum_{i=1}^{10} (x_i 5)^2$. How close is the estimate of σ^2 to the true variance of 9?
- (c) Using your estimate m of the mean, estimate σ^2 by the formula $\sigma^2 = \frac{1}{10} \sum_{i=1}^{10} (x_i m)^2$. How close is the estimate of σ^2 to the true variance of 9?
- (d) Using your estimate m of the mean, estimate σ^2 by the formula $\sigma^2 = \frac{1}{9} \sum_{i=1}^{10} (x_i m)^2$. How close is the estimate of σ^2 to the true variance of 9?

Exercise 2.51 Create a list of the five most important things that you learned about high dimensions.

Exercise 2.52 Write a short essay whose purpose is to excite a college freshman to learn about high dimensions.

3 Best-Fit Subspaces and Singular Value Decomposition (SVD)

3.1 Introduction and Overview

In this chapter, we examine the Singular Value Decomposition (SVD) of a matrix. Consider each row of an $n \times d$ matrix A as a point in d-dimensional space. The singular value decomposition finds the best-fitting k-dimensional subspace for $k = 1, 2, 3, \ldots$, for the set of n data points. Here, "best" means minimizing the sum of the squares of the perpendicular distances of the points to the subspace, or equivalently, maximizing the sum of squares of the lengths of the projections of the points onto this subspace.³ We begin with a special case where the subspace is 1-dimensional, namely a line through the origin. We then show that the best-fitting k-dimensional subspace can be found by kapplications of the best fitting line algorithm, where on the i^{th} iteration we find the best fit line perpendicular to the previous i - 1 lines. When k reaches the rank of the matrix, from these operations we get an exact decomposition of the matrix called the Singular Value Decomposition.

In matrix notation, the singular value decomposition of a matrix A with real entries (we assume all our matrices have real entries) is the factorization of A into the product of three matrices, $A = UDV^T$, where the columns of U and V are orthonormal⁴ and the matrix D is diagonal with positive real entries. The columns of V are the unit length vectors defining the best fitting lines described above (the i^{th} column being the unit-length vector in the direction of the i^{th} line). The coordinates of a row of U will be the fractions of the corresponding row of A along the direction of each of the lines.

The SVD is useful in many tasks. Often a data matrix A is close to a low rank matrix and it is useful to find a good low rank approximation to A. For any k, the singular value decomposition of A gives the best rank-k approximation to A in a well-defined sense.

If $\mathbf{u_i}$ and $\mathbf{v_i}$ are columns of U and V respectively, then the matrix equation $A = UDV^T$ can be rewritten as

$$A = \sum_{i} d_{ii} \mathbf{u}_{i} \mathbf{v}_{i}^{T}.$$

Since $\mathbf{u}_{\mathbf{i}}$ is a $n \times 1$ matrix and $\mathbf{v}_{\mathbf{i}}$ is a $d \times 1$ matrix, $\mathbf{u}_{\mathbf{i}} \mathbf{v}_{\mathbf{i}}^T$ is an $n \times d$ matrix with the same dimensions as A. The i^{th} term in the above sum can be viewed as giving the components of the rows of A along direction $\mathbf{v}_{\mathbf{i}}$. When the terms are summed, they reconstruct A.

³This equivalence is due to the Pythagorean Theorem. For each point, its squared length (its distance to the origin squared) is exactly equal to the squared length of its projection onto the subspace plus the squared distance of the point to its projection; therefore, maximizing the sum of the former is equivalent to minimizing the sum of the latter. For further discussion see Section 3.2.

⁴A set of vectors is orthonormal if each is of length one and they are pairwise orthogonal.

This decomposition of A can be viewed as analogous to writing a vector \mathbf{x} in some orthonormal basis $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_d}$. The coordinates of $\mathbf{x} = (\mathbf{x} \cdot \mathbf{v_1}, \mathbf{x} \cdot \mathbf{v_2} \ldots, \mathbf{x} \cdot \mathbf{v_d})$ are the projections of \mathbf{x} onto the $\mathbf{v_i}$'s. For SVD, this basis has the property that for any k, the first k vectors of this basis produce the least possible total sum of squares error for that value of k.

In addition to the singular value decomposition, there is an eigenvalue decomposition. Let A be a square matrix. A vector \mathbf{v} such that $A\mathbf{v} = \lambda \mathbf{v}$ is called an eigenvector and λ the eigenvalue. When A satisfies a few additional conditions besides being square, the eigenvectors are orthogonal and A can be expressed as $A = VDV^T$ where the eigenvectors are the columns of V and D is a diagonal matrix with the corresponding eigenvalues on its diagonal. If A is symmetric and has distinct singular values, then the singular vectors of A are the eigenvectors. If a singular value has multiplicity d greater than one, the corresponding singular vectors span a subspace of dimension d and any orthogonal basis of the subspace can be used as the eigenvectors or singular vectors. ⁵

The singular value decomposition is defined for all matrices, whereas the more familiar eigenvector decomposition requires that the matrix A be square and certain other conditions on the matrix to ensure orthogonality of the eigenvectors. In contrast, the columns of V in the singular value decomposition, called the *right-singular vectors* of A, always form an orthogonal set with no assumptions on A. The columns of U are called the *left-singular vectors* and they also form an orthogonal set (see Section 3.6). A simple consequence of the orthonormality is that for a square and invertible matrix A, the inverse of A is $VD^{-1}U^{T}$.

Eigenvalues and eignevectors satisfy $A\mathbf{v} = \lambda \mathbf{v}$. We will show that singular values and vectors satisfy a somewhat analogous relationship. Since $A\mathbf{v_i}$ is a $n \times 1$ matrix (vector), the matrix A cannot act on it from the left. But A^T , which is a $d \times n$ matrix, can act on this vector. Indeed, we will show that

$$A\mathbf{v}_{\mathbf{i}} = d_{ii}\mathbf{u}_{\mathbf{i}}$$
 and $A^T\mathbf{u}_{\mathbf{i}} = d_{ii}\mathbf{v}_{\mathbf{i}}$.

In words, A acting on $\mathbf{v_i}$ produces a scalar multiple of $\mathbf{u_i}$ and A^T acting on $\mathbf{u_i}$ produces the same scalar multiple of $\mathbf{v_i}$. Note that $A^T A \mathbf{v_i} = d_{ii}^2 \mathbf{v_i}$. The i^{th} singular vector of A is the i^{th} eigenvector of the square symmetric matrix $A^T A$.

3.2 Preliminaries

Consider projecting a point $\mathbf{a}_{\mathbf{i}} = (a_{i1}, a_{i2}, \dots, a_{id})$ onto a line through the origin. Then

 $a_{i1}^2 + a_{i2}^2 + \dots + a_{id}^2 = (\text{length of projection})^2 + (\text{distance of point to line})^2.$

⁵When d = 1 there are actually two possible singular vectors, one the negative of the other. The subspace spanned is unique.

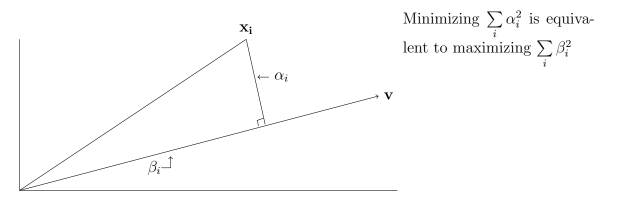


Figure 3.1: The projection of the point \mathbf{x}_i onto the line through the origin in the direction of \mathbf{v} .

This holds by the Pythagorean Theorem (see Figure 3.1). Thus

(distance of point to line)² = $a_{i1}^2 + a_{i2}^2 + \dots + a_{id}^2 - (\text{length of projection})^2$.

Since $\sum_{i=1}^{n} (a_{i1}^2 + a_{i2}^2 + \cdots + a_{id}^2)$ is a constant independent of the line, minimizing the sum of the squares of the distances to the line is equivalent to maximizing the sum of the squares of the lengths of the projections onto the line. Similarly for best-fit subspaces, maximizing the sum of the squared lengths of the projections onto the subspace minimizes the sum of squared distances to the subspace.

Thus we have two interpretations of the best-fit subspace. The first is that it minimizes the sum of squared distances of the data points to it. This interpretation and its use are akin to the notion of least-squares fit from calculus.⁶ The second interpretation of best-fit-subspace is that it maximizes the sum of projections squared of the data points on it. This says that the subspace contains the maximum content of data among all subspaces of the same dimension.

The reader may wonder why we minimize the sum of squared perpendicular distances to the line rather than, say, the sum of distances (not squared). There are examples where the latter definition gives a different answer than the line minimizing the sum of squared perpendicular distances. The choice of the objective function as the sum of squared distances seems a bit arbitrary and in a way it is. But the square has many nice mathematical properties. The first of these, as we have just seen, is that minimizing the sum of squared distances is equivalent to maximizing the sum of squared projections.

⁶But there is a difference: here we take the perpendicular distance to the line or subspace, whereas, in the calculus notion, given *n* pairs, $(x_1, y_1), (x_2, y_2), \ldots, (x_n, y_n)$, we find a line $l = \{(x, y) | y = mx + b\}$ minimizing the vertical squared distances of the points to it, namely, $\sum_{i=1}^{n} (y_i - mx_i - b)^2$.

3.3 Singular Vectors

We now define the *singular vectors* of an $n \times d$ matrix A. Consider the rows of A as n points in a d-dimensional space. Consider the best fit line through the origin. Let \mathbf{v} be a unit vector along this line. The length of the projection of $\mathbf{a_i}$, the i^{th} row of A, onto \mathbf{v} is $|\mathbf{a_i} \cdot \mathbf{v}|$. From this we see that the sum of the squared lengths of the projections is $|A\mathbf{v}|^2$. The best fit line is the one maximizing $|A\mathbf{v}|^2$ and hence minimizing the sum of the squared distances of the points to the line.

With this in mind, define the first singular vector $\mathbf{v_1}$ of A, a column vector, as

$$\mathbf{v_1} = \arg \max_{|\mathbf{v}|=1} |A\mathbf{v}|.$$

Technically, there may be a tie for the vector attaining the maximum and so we should not use the article "the"; in fact, $-\mathbf{v_1}$ is always as good as $\mathbf{v_1}$. In this case, we arbitrarily pick one of the vectors achieving the maximum and refer to it as "the first singular vector" avoiding the more cumbersome "one of the vectors achieving the maximum". We adopt this terminology for all uses of arg max.

The value $\sigma_1(A) = |A\mathbf{v}_1|$ is called the *first singular value* of A. Note that $\sigma_1^2 = \sum_{i=1}^n (\mathbf{a}_i \cdot \mathbf{v}_1)^2$ is the sum of the squared lengths of the projections of the points onto the line determined by \mathbf{v}_1 .

If the data points were all either on a line or close to a line, intuitively, $\mathbf{v_1}$ should give us the direction of that line. It is possible that data points are not close to one line, but lie close to a 2-dimensional subspace or more generally a low dimensional space. Suppose we have an algorithm for finding $\mathbf{v_1}$ (we will describe one such algorithm later). How do we use this to find the best-fit 2-dimensional plane or more generally the best fit *k*-dimensional space?

The greedy approach begins by finding $\mathbf{v_1}$ and then finds the best 2-dimensional subspace containing $\mathbf{v_1}$. The sum of squared distances helps. For every 2-dimensional subspace containing $\mathbf{v_1}$, the sum of squared lengths of the projections onto the subspace equals the sum of squared projections onto $\mathbf{v_1}$ plus the sum of squared projections along a vector perpendicular to $\mathbf{v_1}$ in the subspace. Thus, instead of looking for the best 2dimensional subspace containing $\mathbf{v_1}$, look for a unit vector $\mathbf{v_2}$ perpendicular to $\mathbf{v_1}$ that maximizes $|A\mathbf{v}|^2$ among all such unit vectors. Using the same greedy strategy to find the best three and higher dimensional subspaces, defines $\mathbf{v_3}, \mathbf{v_4}, \ldots$ in a similar manner. This is captured in the following definitions. There is no apriori guarantee that the greedy algorithm gives the best fit. But, in fact, the greedy algorithm does work and yields the best-fit subspaces of every dimension as we will show.

The second singular vector, \mathbf{v}_2 , is defined by the best fit line perpendicular to \mathbf{v}_1 .

$$\mathbf{v_2} = \underset{\substack{\mathbf{v} \perp \mathbf{v_1} \\ |\mathbf{v}|=1}}{\operatorname{arg\,max}} |A\mathbf{v}|$$

The value $\sigma_2(A) = |A\mathbf{v}_2|$ is called the second singular value of A. The third singular vector \mathbf{v}_3 and third singular value are defined similarly by

$$\mathbf{v_3} = \operatorname*{arg\,max}_{\substack{\mathbf{v} \perp \mathbf{v_1}, \mathbf{v_2} \\ |\mathbf{v}|=1}} |A\mathbf{v}|$$

and

$$\sigma_3(A) = |A\mathbf{v}_3|,$$

and so on. The process stops when we have found singular vectors $\mathbf{v}_1, \mathbf{v}_2, \ldots, \mathbf{v}_r$, singular values $\sigma_1, \sigma_2, \ldots, \sigma_r$, and

$$\max_{\substack{\mathbf{v}\perp\mathbf{v_1},\mathbf{v_2},\ldots,\mathbf{v_r}\\ |\mathbf{v}|=1}} |A\mathbf{v}| = 0.$$

The greedy algorithm found the \mathbf{v}_1 that maximized $|A\mathbf{v}|$ and then the best fit 2dimensional subspace containing \mathbf{v}_1 . Is this necessarily the best-fit 2-dimensional subspace overall? The following theorem establishes that the greedy algorithm finds the best subspaces of every dimension.

Theorem 3.1 (The Greedy Algorithm Works) Let A be an $n \times d$ matrix with singular vectors $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_r}$. For $1 \le k \le r$, let V_k be the subspace spanned by $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_k}$. For each k, V_k is the best-fit k-dimensional subspace for A.

Proof: The statement is obviously true for k = 1. For k = 2, let W be a best-fit 2dimensional subspace for A. For any orthonormal basis $(\mathbf{w_1}, \mathbf{w_2})$ of W, $|A\mathbf{w_1}|^2 + |A\mathbf{w_2}|^2$ is the sum of squared lengths of the projections of the rows of A onto W. Choose an orthonormal basis $(\mathbf{w_1}, \mathbf{w_2})$ of W so that $\mathbf{w_2}$ is perpendicular to $\mathbf{v_1}$. If $\mathbf{v_1}$ is perpendicular to W, any unit vector in W will do as $\mathbf{w_2}$. If not, choose $\mathbf{w_2}$ to be the unit vector in Wperpendicular to the projection of $\mathbf{v_1}$ onto W. This makes $\mathbf{w_2}$ perpendicular to $\mathbf{v_1}$.⁷ Since $\mathbf{v_1}$ maximizes $|A\mathbf{v}|^2$, it follows that $|A\mathbf{w_1}|^2 \leq |A\mathbf{v_1}|^2$. Since $\mathbf{v_2}$ maximizes $|A\mathbf{v}|^2$ over all \mathbf{v} perpendicular to $\mathbf{v_1}$, $|A\mathbf{w_2}|^2 \leq |A\mathbf{v_2}|^2$. Thus

$$|A\mathbf{w_1}|^2 + |A\mathbf{w_2}|^2 \le |A\mathbf{v_1}|^2 + |A\mathbf{v_2}|^2.$$

Hence, V_2 is at least as good as W and so is a best-fit 2-dimensional subspace.

For general k, proceed by induction. By the induction hypothesis, V_{k-1} is a best-fit k-1 dimensional subspace. Suppose W is a best-fit k-dimensional subspace. Choose an

⁷This can be seen by noting that $\mathbf{v_1}$ is the sum of two vectors that each are individually perpendicular to $\mathbf{w_2}$, namely the projection of $\mathbf{v_1}$ to W and the portion of $\mathbf{v_1}$ orthogonal to W.

orthonormal basis $\mathbf{w_1}, \mathbf{w_2}, \ldots, \mathbf{w_k}$ of W so that $\mathbf{w_k}$ is perpendicular to $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_{k-1}}$. Then

$$|A\mathbf{w}_{1}|^{2} + |A\mathbf{w}_{2}|^{2} + \dots + |A\mathbf{w}_{k-1}|^{2} \le |A\mathbf{v}_{1}|^{2} + |A\mathbf{v}_{2}|^{2} + \dots + |A\mathbf{v}_{k-1}|^{2}$$

since V_{k-1} is an optimal k-1 dimensional subspace. Since $\mathbf{w}_{\mathbf{k}}$ is perpendicular to $\mathbf{v}_{1}, \mathbf{v}_{2}, \ldots, \mathbf{v}_{\mathbf{k}-1}$, by the definition of $\mathbf{v}_{\mathbf{k}}, |A\mathbf{w}_{\mathbf{k}}|^{2} \leq |A\mathbf{v}_{\mathbf{k}}|^{2}$. Thus $|A\mathbf{w}_{1}|^{2} + |A\mathbf{w}_{2}|^{2} + \cdots + |A\mathbf{w}_{\mathbf{k}-1}|^{2} + |A\mathbf{w}_{\mathbf{k}}|^{2} \leq |A\mathbf{v}_{1}|^{2} + |A\mathbf{v}_{2}|^{2} + \cdots + |A\mathbf{v}_{\mathbf{k}-1}|^{2} + |A\mathbf{v}_{\mathbf{k}}|^{2}$, $|A\mathbf{w}_{\mathbf{k}}|^{2} \leq |A\mathbf{v}_{1}|^{2} + |A\mathbf{v}_{2}|^{2} + \cdots + |A\mathbf{v}_{\mathbf{k}-1}|^{2} + |A\mathbf{v}_{\mathbf{k}}|^{2}$, $|A\mathbf{v}_{\mathbf{k}}|^{2} \leq |A\mathbf{v}_{\mathbf{k}}|^{2}$.

proving that V_k is at least as good as W and hence is optimal.

Note that the *n*-dimensional vector $A\mathbf{v_i}$ is a list of lengths (with signs) of the projections of the rows of A onto $\mathbf{v_i}$. Think of $|A\mathbf{v_i}| = \sigma_i(A)$ as the "component" of the matrix A along $\mathbf{v_i}$. For this interpretation to make sense, it should be true that adding up the squares of the components of A along each of the $\mathbf{v_i}$ gives the square of the "whole content of the matrix A". This is indeed the case and is the matrix analogy of decomposing a vector into its components along orthogonal directions.

Consider one row, say $\mathbf{a}_{\mathbf{j}}$, of A. Since $\mathbf{v}_1, \mathbf{v}_2, \dots, \mathbf{v}_{\mathbf{r}}$ span the space of all rows of A, $\mathbf{a}_{\mathbf{j}} \cdot \mathbf{v} = 0$ for all \mathbf{v} perpendicular to $\mathbf{v}_1, \mathbf{v}_2, \dots, \mathbf{v}_{\mathbf{r}}$. Thus, for each row $\mathbf{a}_{\mathbf{j}}, \sum_{i=1}^r (\mathbf{a}_{\mathbf{j}} \cdot \mathbf{v}_i)^2 = |\mathbf{a}_{\mathbf{j}}|^2$. Summing over all rows j,

$$\sum_{j=1}^{n} |\mathbf{a}_{j}|^{2} = \sum_{j=1}^{n} \sum_{i=1}^{r} (\mathbf{a}_{j} \cdot \mathbf{v}_{i})^{2} = \sum_{i=1}^{r} \sum_{j=1}^{n} (\mathbf{a}_{j} \cdot \mathbf{v}_{i})^{2} = \sum_{i=1}^{r} |A\mathbf{v}_{i}|^{2} = \sum_{i=1}^{r} \sigma_{i}^{2}(A).$$

But $\sum_{j=1}^{n} |\mathbf{a}_{j}|^{2} = \sum_{j=1}^{n} \sum_{k=1}^{d} a_{jk}^{2}$, the sum of squares of all the entries of A. Thus, the sum of squares of the singular values of A is indeed the square of the "whole content of A", i.e., the sum of squares of all the entries. There is an important norm associated with this quantity, the Frobenius norm of A, denoted $||A||_{F}$ defined as

$$||A||_F = \sqrt{\sum_{j,k} a_{jk}^2}.$$

Lemma 3.2 For any matrix A, the sum of squares of the singular values equals the square of the Frobenius norm. That is, $\sum \sigma_i^2(A) = ||A||_F^2$.

Proof: By the preceding discussion.

The vectors $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_r}$ are called the *right-singular vectors*. The vectors $A\mathbf{v_i}$ form a fundamental set of vectors and we normalize them to length one by

$$\mathbf{u}_{\mathbf{i}} = \frac{1}{\sigma_i(A)} A \mathbf{v}_{\mathbf{i}}.$$

Later we will show that $\mathbf{u}_1, \mathbf{u}_2, \ldots, \mathbf{u}_r$ similarly maximize $|\mathbf{u}^T A|$ when multiplied on the left and are called the *left-singular vectors*. Clearly, the right-singular vectors are orthogonal by definition. We will show later that the left-singular vectors are also orthogonal.

3.4 Singular Value Decomposition (SVD)

Let A be an $n \times d$ matrix with singular vectors $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_r}$ and corresponding singular values $\sigma_1, \sigma_2, \ldots, \sigma_r$. The left-singular vectors of A are $\mathbf{u_i} = \frac{1}{\sigma_i} A \mathbf{v_i}$ where $\sigma_i \mathbf{u_i}$ is a vector whose coordinates correspond to the projections of the rows of A onto $\mathbf{v_i}$. Each $\sigma_i \mathbf{u_i v_i^T}$ is a rank one matrix whose rows are the " $\mathbf{v_i}$ components" of the rows of A, i.e., the projections of the rows of A in the $\mathbf{v_i}$ direction. We will prove that A can be decomposed into a sum of rank one matrices as

$$A = \sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T.$$

Geometrically, each point is decomposed in A into its components along each of the r orthogonal directions given by the \mathbf{v}_i . We will also prove this algebraically. We begin with a simple lemma that two matrices A and B are identical if $A\mathbf{v} = B\mathbf{v}$ for all \mathbf{v} .

Lemma 3.3 Matrices A and B are identical if and only if for all vectors \mathbf{v} , $A\mathbf{v} = B\mathbf{v}$.

Proof: Clearly, if A = B then $A\mathbf{v} = B\mathbf{v}$ for all \mathbf{v} . For the converse, suppose that $A\mathbf{v} = B\mathbf{v}$ for all \mathbf{v} . Let \mathbf{e}_i be the vector that is all zeros except for the i^{th} component which has value one. Now $A\mathbf{e}_i$ is the i^{th} column of A and thus A = B if for each i, $A\mathbf{e}_i = B\mathbf{e}_i$.

Theorem 3.4 Let A be an $n \times d$ matrix with right-singular vectors $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_r}$, leftsingular vectors $\mathbf{u_1}, \mathbf{u_2}, \ldots, \mathbf{u_r}$, and corresponding singular values $\sigma_1, \sigma_2, \ldots, \sigma_r$. Then

$$A = \sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T.$$

Proof: We first show that multiplying A and $\sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ by \mathbf{v}_j results in equality.

$$\sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T \mathbf{v}_j = \sigma_j \mathbf{u}_j = A \mathbf{v}_j$$

Since any vector \mathbf{v} can be expressed as a linear combination of the singular vectors plus a vector perpendicular to the \mathbf{v}_i , $A\mathbf{v} = \sum_{i=1}^r \sigma_i \mathbf{u}_i \mathbf{v}_i^T \mathbf{v}$ for all \mathbf{v} and by Lemma 3.3, $A = \sum_{i=1}^r \sigma_i \mathbf{u}_i \mathbf{v}_i^T$.

The decomposition $A = \sum_i \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ is called the *singular value decomposition*, *SVD*, of A. In matrix notation $A = UDV^T$ where the columns of U and V consist of the left and right-singular vectors, respectively, and D is a diagonal matrix whose diagonal entries are

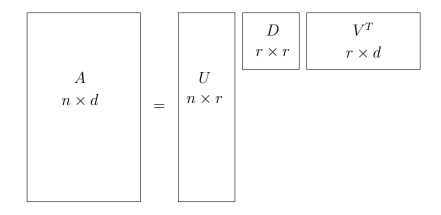


Figure 3.2: The SVD decomposition of an $n \times d$ matrix.

the singular values of A. For any matrix A, the sequence of singular values is unique and if the singular values are all distinct, then the sequence of singular vectors is unique up to signs. However, when some set of singular values are equal, the corresponding singular vectors span some subspace. Any set of orthonormal vectors spanning this subspace can be used as the singular vectors.

3.5 Best Rank-*k* Approximations

Let A be an $n \times d$ matrix and think of the rows of A as n points in d-dimensional space. Let

$$A = \sum_{i=1}^{r} \sigma_i \mathbf{u_i v_i^T}$$

be the SVD of A. For $k \in \{1, 2, \dots, r\}$, let

$$A_k = \sum_{i=1}^k \sigma_i \mathbf{u}_i \mathbf{v}_i^T$$

be the sum truncated after k terms. It is clear that A_k has rank k. We show that A_k is the best rank k approximation to A, where, error is measured in the Frobenius norm. Geometrically, this says that $\mathbf{v_1}, \ldots, \mathbf{v_k}$ define the k-dimensional space minimizing the sum of squared distances of the points to the space. To see why, we need the following lemma.

Lemma 3.5 The rows of A_k are the projections of the rows of A onto the subspace V_k spanned by the first k singular vectors of A.

Proof: Let **a** be an arbitrary row vector. Since the **v**_i are orthonormal, the projection of the vector **a** onto V_k is given by $\sum_{i=1}^{k} (\mathbf{a} \cdot \mathbf{v}_i) \mathbf{v}_i^T$. Thus, the matrix whose rows are

the projections of the rows of A onto V_k is given by $\sum_{i=1}^k A \mathbf{v}_i \mathbf{v}_i^T$. This last expression simplifies to

$$\sum_{i=1}^{k} A \mathbf{v}_{i} \mathbf{v}_{i}^{T} = \sum_{i=1}^{k} \sigma_{i} \mathbf{u}_{i} \mathbf{v}_{i}^{T} = A_{k}.$$

Theorem 3.6 For any matrix B of rank at most k

$$||A - A_k||_F \le ||A - B||_F$$

Proof: Let *B* minimize $||A - B||_F^2$ among all rank *k* or less matrices. Let *V* be the space spanned by the rows of *B*. The dimension of *V* is at most *k*. Since *B* minimizes $||A - B||_F^2$, it must be that each row of *B* is the projection of the corresponding row of *A* onto *V*: Otherwise replace the row of *B* with the projection of the corresponding row of *A* onto *V*. This still keeps the row space of *B* contained in *V* and hence the rank of *B* is still at most *k*. But it reduces $||A - B||_F^2$, contradicting the minimality of $||A - B||_F$.

Since each row of B is the projection of the corresponding row of A, it follows that $||A - B||_F^2$ is the sum of squared distances of rows of A to V. Since A_k minimizes the sum of squared distance of rows of A to any k-dimensional subspace, from Theorem 3.1, it follows that $||A - A_k||_F \leq ||A - B||_F$.

In addition to Frobenius norm, there is another matrix norm of interest. To motivate, consider the example of a term-document matrix A. Suppose we have a large database of documents that form rows of an $n \times d$ matrix A. There are d terms and each document is a d-dimensional vector with one component per term, which is the number of occurrences of the term in the document. We are allowed to "preprocess" A. After the preprocessing, we receive queries. Each query \mathbf{x} is an d-dimensional vector which specifies how important each term is to the query. The desired answer is a *n*-dimensional vector which gives the similarity (dot product) of the query to each document in the database, namely $A\mathbf{x}$, the "matrix-vector" product. Query time is to be much less than preprocessing time, since the idea is that we need to answer many queries for the same database. Besides this, there are many situations where one performs many matrix vector products with the same matrix. This is applicable to these situations as well. Näively, it would take O(nd) time to do the product $A\mathbf{x}$. Suppose we computed the SVD and took $A_k = \sum_{i=1}^k \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ as our approximation to A. Then, we could return $A_k \mathbf{x} = \sum_{i=1}^k \sigma_i \mathbf{u}_i (\mathbf{v}_i \cdot \mathbf{x})$ as the approximation to $A\mathbf{x}$. This only takes k dot products of d-dimensional vectors, followed by a sum of k ndimensional vectors, and so takes time O(kd+kn), which is a win provided $k \ll \min(d, n)$. How is the error measured? Since \mathbf{x} is unknown, the approximation needs to be good for every **x**. So we should take the maximum over all **x** of $|(A_k - A)\mathbf{x}|$. This would be infinite since $|\mathbf{x}|$ can grow without bound. So we restrict to $|\mathbf{x}| \leq 1$. Formally, we define a new norm of a matrix A by

$$||A||_2 = \max_{\mathbf{x}:|\mathbf{x}| \le 1} |A\mathbf{x}|.$$

This is called the 2-norm or the spectral norm. Note that it equals $\sigma_1(A)$.

3.6 Left Singular Vectors

Theorem 3.7 The left singular vectors are pairwise orthogonal.

Proof: First we show that each $\mathbf{u}_i, i \geq 2$ is orthogonal to \mathbf{u}_1 . Suppose not, and for some $i \geq 2$, $\mathbf{u}_1^T \mathbf{u}_i \neq 0$. Without loss of generality assume that $\mathbf{u}_1^T \mathbf{u}_i = \delta > 0$. (If $\mathbf{u}_1^T \mathbf{u}_i < 0$ then just replace \mathbf{u}_i with $-\mathbf{u}_i$.) For $\varepsilon > 0$, let

$$\mathbf{v}_1' = \frac{\mathbf{v}_1 + \varepsilon \mathbf{v}_i}{|\mathbf{v}_1 + \varepsilon \mathbf{v}_i|}.$$

Notice that \mathbf{v}'_1 is a unit-length vector.

$$A\mathbf{v}_1' = \frac{\sigma_1 \mathbf{u}_1 + \varepsilon \sigma_i \mathbf{u}_i}{\sqrt{1 + \varepsilon^2}}$$

has length at least as large as its component along \mathbf{u}_1 which is

$$\mathbf{u_1^T}(\frac{\sigma_1\mathbf{u_1} + \varepsilon\sigma_i\mathbf{u_i}}{\sqrt{1 + \varepsilon^2}}) > (\sigma_1 + \varepsilon\sigma_i\delta)\left(1 - \frac{\varepsilon^2}{2}\right) > \sigma_1 - \frac{\varepsilon^2}{2}\sigma_1 + \varepsilon\sigma_i\delta + \frac{\varepsilon^3}{2}\sigma_i\delta > \sigma_1,$$

for sufficiently small ϵ , a contradiction to the definition of σ_1 . Thus $\mathbf{u}_1 \cdot \mathbf{u}_i = 0$ for $i \geq 2$.

The proof for other \mathbf{u}_i and \mathbf{u}_j , j > i > 1 is similar. Suppose without loss of generality that $\mathbf{u}_i^T \mathbf{u}_j > \delta > 0$.

$$A\left(\frac{\mathbf{v}_i + \varepsilon \mathbf{v}_j}{|\mathbf{v}_i + \varepsilon \mathbf{v}_j|}\right) = \frac{\sigma_i \mathbf{u}_i + \varepsilon \sigma_j \mathbf{u}_j}{\sqrt{1 + \varepsilon^2}}$$

has length at least as large as its component along \mathbf{u}_i which is

$$\mathbf{u_i^T}(\frac{\sigma_1\mathbf{u_i} + \varepsilon\sigma_j\mathbf{u_j}}{\sqrt{1+\varepsilon^2}}) > \left(\sigma_i + \varepsilon\sigma_j\mathbf{u_i^T}\mathbf{u_j}\right) \left(1 - \frac{\varepsilon^2}{2}\right) > \sigma_i - \frac{\varepsilon^2}{2}\sigma_i + \varepsilon\sigma_j\delta - \frac{\varepsilon^3}{2}\sigma_i\delta > \sigma_i,$$

for sufficiently small ϵ , a contradiction since $\mathbf{v_i} + \varepsilon \mathbf{v_j}$ is orthogonal to $\mathbf{v_1}, \mathbf{v_2}, \dots, \mathbf{v_{i-1}}$ and σ_i is defined to be the maximum of $|A\mathbf{v}|$ over such vectors.

Next we prove that A_k is the best rank k, 2-norm approximation to A. We first show that the square of the 2-norm of $A - A_k$ is the square of the $(k+1)^{st}$ singular value of A. This is essentially by definition of A_k ; that is, A_k represents the projections of the points in A onto the space spanned by the top k singular vectors, and so $A - A_k$ is the remaining portion of those points, whose top singular value will be σ_{k+1} .

Lemma 3.8 $||A - A_k||_2^2 = \sigma_{k+1}^2$.

Proof: Let $A = \sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ be the singular value decomposition of A. Then $A_k = \sum_{i=1}^{k} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ and $A - A_k = \sum_{i=k+1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$. Let \mathbf{v} be the top singular vector of $A - A_k$. Express \mathbf{v} as a linear combination of $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_r}$. That is, write $\mathbf{v} = \sum_{i=1}^r c_i \mathbf{v_i}$. Then

$$|(A - A_k)\mathbf{v}| = \left| \sum_{i=k+1}^r \sigma_i \mathbf{u}_i \mathbf{v}_i^T \sum_{j=1}^r c_j \mathbf{v}_j \right| = \left| \sum_{i=k+1}^r c_i \sigma_i \mathbf{u}_i \mathbf{v}_i^T \mathbf{v}_i \right|$$
$$= \left| \sum_{i=k+1}^r c_i \sigma_i \mathbf{u}_i \right| = \sqrt{\sum_{i=k+1}^r c_i^2 \sigma_i^2},$$

since the \mathbf{u}_i are orthonormal. The \mathbf{v} maximizing this last quantity, subject to the constraint that $|\mathbf{v}|^2 = \sum_{i=1}^r c_i^2 = 1$, occurs when $c_{k+1} = 1$ and the rest of the c_i are zero. Thus, $||A - A_k||_2^2 = \sigma_{k+1}^2$ proving the lemma.

Finally, we prove that A_k is the best rank k, 2-norm approximation to A:

Theorem 3.9 Let A be an $n \times d$ matrix. For any matrix B of rank at most k

$$||A - A_k||_2 \le ||A - B||_2.$$

Proof: If A is of rank k or less, the theorem is obviously true since $||A - A_k||_2 = 0$. Assume that A is of rank greater than k. By Lemma 3.8, $||A - A_k||_2^2 = \sigma_{k+1}^2$. The null space of B, the set of vectors **v** such that $B\mathbf{v} = 0$, has dimension at least d - k. Let $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_{k+1}}$ be the first k + 1 singular vectors of A. By a dimension argument, it follows that there exists a $\mathbf{z} \neq 0$ in

Null
$$(B) \cap$$
 Span $\{\mathbf{v_1}, \mathbf{v_2}, \dots, \mathbf{v_{k+1}}\}$.

Scale \mathbf{z} to be of length one.

$$||A - B||_{2}^{2} \ge |(A - B)\mathbf{z}|^{2}.$$

Since $B\mathbf{z} = 0$,

$$||A - B||_2^2 \ge |A\mathbf{z}|^2$$
.

Since \mathbf{z} is in the $\mathrm{Span}\left\{\mathbf{v_1},\mathbf{v_2},\ldots,\mathbf{v_{k+1}}\right\}$

$$|A\mathbf{z}|^{2} = \left|\sum_{i=1}^{n} \sigma_{i} \mathbf{u}_{i} \mathbf{v}_{i}^{T} \mathbf{z}\right|^{2} = \sum_{i=1}^{n} \sigma_{i}^{2} \left(\mathbf{v}_{i}^{T} \mathbf{z}\right)^{2} = \sum_{i=1}^{k+1} \sigma_{i}^{2} \left(\mathbf{v}_{i}^{T} \mathbf{z}\right)^{2} \ge \sigma_{k+1}^{2} \sum_{i=1}^{k+1} \left(\mathbf{v}_{i}^{T} \mathbf{z}\right)^{2} = \sigma_{k+1}^{2}.$$

It follows that $||A - B||_2^2 \ge \sigma_{k+1}^2$ proving the theorem.

We now prove the analog of eigenvectors and eigenvalues for singular values and vectors we discussed in the introduction.

Lemma 3.10 (Analog of eigenvalues and eigenvectors)

$$A\mathbf{v}_{\mathbf{i}} = \sigma_i \mathbf{u}_{\mathbf{i}} \text{ and } A^T \mathbf{u}_{\mathbf{i}} = \sigma_i \mathbf{v}_{\mathbf{i}}.$$

Proof: The first equation is already known. For the second, note that from the SVD, we get $A^T \mathbf{u_i} = \sum_j \sigma_j \mathbf{v_j} \mathbf{u_j}^T \mathbf{u_i}$, where since the $\mathbf{u_j}$ are orthonormal, all terms in the summation are zero except for j = i.

3.7 Power Method for Computing the Singular Value Decomposition

Computing the singular value decomposition is an important branch of numerical analysis in which there have been many sophisticated developments over a long period of time. The reader is referred to numerical analysis texts for more details. Here we present an "in-principle" method to establish that the approximate SVD of a matrix A can be computed in polynomial time. The method we present, called the *power method*, is simple and is in fact the conceptual starting point for many algorithms. Let A be a matrix whose SVD is $\sum_{i} \sigma_{i} \mathbf{u}_{i} \mathbf{v}_{i}^{T}$. We wish to work with a matrix that is square and symmetric. Let $B = A^{T}A$. By direct multiplication, using the orthogonality of the \mathbf{u}_{i} 's that was proved in Theorem 3.7,

$$B = A^{T}A = \left(\sum_{i} \sigma_{i} \mathbf{v}_{i} \mathbf{u}_{i}^{T}\right) \left(\sum_{j} \sigma_{j} \mathbf{u}_{j} \mathbf{v}_{j}^{T}\right)$$
$$= \sum_{i,j} \sigma_{i} \sigma_{j} \mathbf{v}_{i} (\mathbf{u}_{i}^{T} \cdot \mathbf{u}_{j}) \mathbf{v}_{j}^{T} = \sum_{i} \sigma_{i}^{2} \mathbf{v}_{i} \mathbf{v}_{i}^{T}.$$

The matrix *B* is square and symmetric, and has the same left and right-singular vectors. In particular, $B\mathbf{v_j} = (\sum_i \sigma_i^2 \mathbf{v_i v_i^T})\mathbf{v_j} = \sigma_j^2 \mathbf{v_j}$, so $\mathbf{v_j}$ is an eigenvector of *B* with eigenvalue σ_j^2 . If *A* is itself square and symmetric, it will have the same right and left-singular vectors, namely $A = \sum_i \sigma_i \mathbf{v_i v_i^T}$ and computing *B* is unnecessary.

Now consider computing B^2 .

$$B^{2} = \left(\sum_{i} \sigma_{i}^{2} \mathbf{v}_{i} \mathbf{v}_{i}^{T}\right) \left(\sum_{j} \sigma_{j}^{2} \mathbf{v}_{j} \mathbf{v}_{j}^{T}\right) = \sum_{ij} \sigma_{i}^{2} \sigma_{j}^{2} \mathbf{v}_{i} (\mathbf{v}_{i}^{T} \mathbf{v}_{j}) \mathbf{v}_{j}^{T}$$

When $i \neq j$, the dot product $\mathbf{v_i}^T \mathbf{v_j}$ is zero by orthogonality.⁸ Thus, $B^2 = \sum_{i=1}^r \sigma_i^4 \mathbf{v_i v_i}^T$. In computing the k^{th} power of B, all the cross product terms are zero and

$$B^k = \sum_{i=1}^r \sigma_i^{2k} \mathbf{v}_i \mathbf{v}_i^T.$$

⁸The "outer product" $\mathbf{v_i v_j}^T$ is a matrix and is not zero even for $i \neq j$.

If $\sigma_1 > \sigma_2$, then the first term in the summation dominates, so $B^k \to \sigma_1^{2k} \mathbf{v_1} \mathbf{v_1}^T$. This means a close estimate to $\mathbf{v_1}$ can be computed by simply taking the first column of B^k and normalizing it to a unit vector.

3.7.1 A Faster Method

A problem with the above method is that A may be a very large, sparse matrix, say a $10^8 \times 10^8$ matrix with 10^9 nonzero entries. Sparse matrices are often represented by just a list of non-zero entries, say, a list of triples of the form (i, j, a_{ij}) . Though A is sparse, B need not be and in the worse case may have all 10^{16} entries non-zero,⁹ and it is then impossible to even write down B, let alone compute the product B^2 . Even if A is moderate in size, computing matrix products is costly in time. Thus, a more efficient method is needed.

Instead of computing B^k , select a random vector \mathbf{x} and compute the product $B^k \mathbf{x}$. The vector \mathbf{x} can be expressed in terms of the singular vectors of B augmented to a full orthonormal basis as $\mathbf{x} = \sum c_i \mathbf{v_i}$. Then

$$B^{k}\mathbf{x} \approx (\sigma_{1}^{2k}\mathbf{v_{1}}\mathbf{v_{1}}^{T}) \left(\sum_{i=1}^{d} c_{i}\mathbf{v_{i}}\right) = \sigma_{1}^{2k}c_{1}\mathbf{v_{1}}.$$

Normalizing the resulting vector yields $\mathbf{v_1}$, the first singular vector of A. The way $B^k \mathbf{x}$ is computed is by a series of matrix vector products, instead of matrix products. $B^k \mathbf{x} = A^T A \dots A^T A \mathbf{x}$, which can be computed right-to-left. This consists of 2k vector times sparse matrix multiplications.

An issue occurs if there is no significant gap between the first and second singular values of a matrix. Take for example the case when there is a tie for the first singular vector and $\sigma_1 = \sigma_2$. Then, the argument above fails. We will overcome this hurdle. Theorem 3.11 below states that even with ties, the power method converges to some vector in the span of those singular vectors corresponding to the "nearly highest" singular values. The theorem needs a vector \mathbf{x} which has a component of at least δ along the first right singular vector \mathbf{v}_1 of A. We will see in Lemma 3.12 that a random vector satisfies this condition.

Theorem 3.11 Let A be an $n \times d$ matrix and \mathbf{x} a unit length vector in \mathbf{R}^d with $|\mathbf{x}^T \mathbf{v_1}| \geq \delta$, where $\delta > 0$. Let V be the space spanned by the right singular vectors of A corresponding to singular values greater than $(1 - \varepsilon) \sigma_1$. Let \mathbf{w} be the unit vector after $k = \frac{\ln(1/\varepsilon\delta)}{2\varepsilon}$ iterations of the power method, namely,

$$\mathbf{w} = \frac{\left(A^{T}A\right)^{k}\mathbf{x}}{\left|\left(A^{T}A\right)^{k}\mathbf{x}\right|}.$$

Then **w** has a component of at most ε perpendicular to V.

 $^{{}^{9}}$ E.g., suppose each entry in the first row of A is non-zero and the rest of A is zero.

Proof: Let

$$A = \sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$$

be the SVD of A. If the rank of A is less than d, then for convenience complete $\{\mathbf{v}_1, \mathbf{v}_2, \ldots \mathbf{v}_r\}$ into an orthonormal basis $\{\mathbf{v}_1, \mathbf{v}_2, \ldots \mathbf{v}_d\}$ of d-space. Write \mathbf{x} in the basis of the \mathbf{v}_i 's as

$$\mathbf{x} = \sum_{i=1}^d c_i \mathbf{v_i}.$$

Since $(A^T A)^k = \sum_{i=1}^d \sigma_i^{2k} \mathbf{v}_i \mathbf{v}_i^T$, it follows that $(A^T A)^k \mathbf{x} = \sum_{i=1}^d \sigma_i^{2k} c_i \mathbf{v}_i$. By hypothesis, $|c_1| \ge \delta$.

Suppose that $\sigma_1, \sigma_2, \ldots, \sigma_m$ are the singular values of A that are greater than or equal to $(1 - \varepsilon) \sigma_1$ and that $\sigma_{m+1}, \ldots, \sigma_d$ are the singular values that are less than $(1 - \varepsilon) \sigma_1$. Now

$$|(A^T A)^k \mathbf{x}|^2 = \left|\sum_{i=1}^d \sigma_i^{2k} c_i \mathbf{v}_i\right|^2 = \sum_{i=1}^d \sigma_i^{4k} c_i^2 \ge \sigma_1^{4k} c_1^2 \ge \sigma_1^{4k} \delta^2.$$

The component of $|(A^T A)^k \mathbf{x}|^2$ perpendicular to the space V is

$$\sum_{i=m+1}^{d} \sigma_i^{4k} c_i^2 \le (1-\varepsilon)^{4k} \sigma_1^{4k} \sum_{i=m+1}^{d} c_i^2 \le (1-\varepsilon)^{4k} \sigma_1^{4k}$$

since $\sum_{i=1}^{d} c_i^2 = |\mathbf{x}| = 1$. Thus, the component of \mathbf{w} perpendicular to V has squared length at most $\frac{(1-\varepsilon)^{4k}\sigma_1^{4k}}{\sigma_1^{4k}\delta^2}$ and so its length is at most

$$\frac{(1-\varepsilon)^{2k}\sigma_1^{2k}}{\delta\sigma_1^{2k}} = \frac{(1-\varepsilon)^{2k}}{\delta} \le \frac{e^{-2k\varepsilon}}{\delta} = \varepsilon.$$

Lemma 3.12 Let $\mathbf{y} \in \mathbf{R}^n$ be a random vector with the unit variance spherical Gaussian as its probability density. Let $\mathbf{x} = \mathbf{y}/|\mathbf{y}|$. Let \mathbf{v} be any fixed (not random) unit length vector. Then

$$Prob\left(|\mathbf{x}^T\mathbf{v}| \le \frac{1}{20\sqrt{d}}\right) \le \frac{1}{10} + 3e^{-d/64}.$$

Proof: With $c = \sqrt{d}$ substituted in Theorem (2.9) of Chapter 2, the probability that $|\mathbf{y}| \geq 2\sqrt{d}$ is at most $3e^{-d/64}$. Further, $\mathbf{y}^T \mathbf{v}$ is a random, zero mean, unit variance Gaussian. Thus, the probability that $|\mathbf{y}^T \mathbf{v}| \leq \frac{1}{10}$ is at most 1/10. Combining these two facts and using the union bound, establishes the lemma.

3.8 Singular Vectors and Eigenvectors

Recall that for a square matrix B, if the vector \mathbf{x} and scalar λ are such that $B\mathbf{x} = \lambda \mathbf{x}$, then \mathbf{x} is an *eigenvector* of B and λ is the corresponding *eigenvalue*. As we saw in Section 3.7, if $B = A^T A$, then the right singular vectors $\mathbf{v_j}$ of A are eigenvectors of B with eigenvalues σ_j^2 . The same argument shows that the left singular vectors $\mathbf{u_j}$ of A are eigenvectors of AA^T with eigenvalues σ_j^2 .

Notice that $B = A^T A$ has the property that for any vector \mathbf{x} we have $\mathbf{x}^T B \mathbf{x} \ge 0$. This is because $B = \sum_i \sigma_i^2 \mathbf{v_i v_i}^T$ and for any \mathbf{x} we have $\mathbf{x}^T \mathbf{v_i v_i}^T \mathbf{x} = (\mathbf{x}^T \mathbf{v_i})^2 \ge 0$. A matrix B with the property that $\mathbf{x}^T B \mathbf{x} \ge 0$ for all \mathbf{x} is called *positive semi-definite*. Every matrix of the form $A^T A$ is positive semi-definite. In the other direction, any positive semi-definite matrix B can be decomposed into a product $A^T A$, and so its eigenvalue decomposition can be obtained from the singular value decomposition of A. The interested reader should consult a linear algebra book.

3.9 Applications of Singular Value Decomposition

3.9.1 Centering Data

Singular value decomposition is used in many applications and for some of these applications it is essential to first center the data by subtracting the centroid of the data from each data point.¹⁰ For instance, if you are interested in the statistics of the data and how it varies in relationship to its mean, then you would want to center the data. On the other hand, if you are interested in finding the best low rank approximation to a matrix, then you do not center the data. The issue is whether you are finding the best fitting subspace or the best fitting affine space; in the latter case you first center the data and then find the best fitting subspace.

We first show that the line minimizing the sum of squared distances to a set of points, if not restricted to go through the origin, must pass through the centroid of the points. This implies that if the centroid is subtracted from each data point, such a line will pass through the origin. The best fit line can be generalized to k dimensional "planes". The operation of subtracting the centroid from all data points is useful in other contexts as well. So we give it the name "centering data".

Lemma 3.13 The best-fit line (minimizing the sum of perpendicular distances squared) of a set of data points must pass through the centroid of the points.

Proof: Subtract the centroid from each data point so that the centroid is **0**. Let ℓ be the best-fit line and assume for contradiction that ℓ does not pass through the origin. The line ℓ can be written as $\{\mathbf{a} + \lambda \mathbf{v} | \lambda \in \mathbf{R}\}$, where **a** is the closest point to **0** on ℓ and **v** is a unit length vector in the direction of ℓ , which is perpendicular to **a**. For a data point

¹⁰The centroid of a set of points is the coordinate-wise average of the points.

 $\mathbf{a_i}$, let $dist(\mathbf{a_i}, \ell)$ denote its perpendicular distance to ℓ . By the Pythagorean theorem, we have $|\mathbf{a_i} - \mathbf{a}|^2 = dist(\mathbf{a_i}, \ell)^2 + (\mathbf{v} \cdot \mathbf{a_i})^2$, or equivalently, $dist(\mathbf{a_i}, \ell)^2 = |\mathbf{a_i} - \mathbf{a}|^2 - (\mathbf{v} \cdot \mathbf{a_i})^2$. Summing over all data points:

$$\begin{split} &\sum_{i=1}^{n} dist(\mathbf{a_{i}}, \ell)^{2} = \sum_{i=1}^{n} \left(|\mathbf{a_{i}} - \mathbf{a}|^{2} - (\mathbf{v} \cdot \mathbf{a_{i}})^{2} \right) = \sum_{i=1}^{n} \left(|\mathbf{a_{i}}|^{2} + |\mathbf{a}|^{2} - 2\mathbf{a_{i}} \cdot \mathbf{a} - (\mathbf{v} \cdot \mathbf{a_{i}})^{2} \right) \\ &= \sum_{i=1}^{n} |\mathbf{a_{i}}|^{2} + n|\mathbf{a}|^{2} - 2\mathbf{a} \cdot \left(\sum_{i} \mathbf{a_{i}} \right) - \sum_{i=1}^{n} (\mathbf{v} \cdot \mathbf{a_{i}})^{2} = \sum_{i} |\mathbf{a_{i}}|^{2} + n|\mathbf{a}|^{2} - \sum_{i} (\mathbf{v} \cdot \mathbf{a_{i}})^{2}, \end{split}$$

where we used the fact that since the centroid is $\mathbf{0}$, $\sum_i \mathbf{a}_i = 0$. The above expression is minimized when $\mathbf{a} = \mathbf{0}$, so the line $\ell' = \{\lambda \mathbf{v} : \lambda \in \mathbf{R}\}$ through the origin is a better fit than ℓ , contradicting ℓ being the best-fit line.

This proof, as well as others, requires the sum of squared distances for the Pythagorean theorem to apply.

A statement analogous to Lemma 3.13 holds for higher dimensional objects. Define an *affine space* as a subspace translated by a vector. So an affine space is a set of the form

$$\{\mathbf{v}_0 + \sum_{i=1}^k c_i \mathbf{v}_i | c_1, c_2, \dots, c_k \in \mathbf{R}\}.$$

Here, $\mathbf{v_0}$ is the translation and $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_k}$ form an orthonormal basis for the subspace.

Lemma 3.14 The k dimensional affine space which minimizes the sum of squared perpendicular distances to the data points must pass through the centroid of the points.

Proof: We only give a brief idea of the proof, which is similar to the previous lemma. Instead of $(\mathbf{v} \cdot \mathbf{a}_i)^2$, we will now have $\sum_{j=1}^k (\mathbf{v}_j \cdot \mathbf{a}_i)^2$, where, \mathbf{v}_j , j = 1, 2, ..., k are an orthonormal basis of the subspace through the origin parallel to the affine space.

3.9.2 Principal Component Analysis

The traditional use of SVD is in Principal Component Analysis (PCA). PCA is illustrated by a movie recommendation setting where there are n customers and d movies. Let matrix A with elements a_{ij} represent the amount that customer i likes movie j. One hypothesizes that there are only k underlying basic factors that determine how much a given customer will like a given movie, where k is much smaller than n or d. For example, these could be the amount of comedy, drama, and action, the novelty of the story, etc. Each movie can be described as a k-dimensional vector indicating how much of these basic factors the movie has, and each customer can be described as a k-dimensional vector indicating how important each of these basic factors is to that customer; the dot-product of these two vectors is hypothesized to determine how much that customer will like that movie. In particular, this means that the $n \times d$ matrix A can be expressed as the product

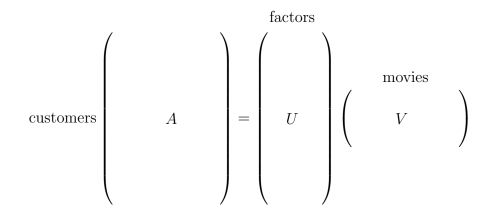


Figure 3.3: Customer-movie data

of an $n \times k$ matrix U describing the customers and a $k \times d$ matrix V describing the movies. Finding the best rank k approximation A_k by SVD gives such a U and V. One twist is that A may not be exactly equal to UV, in which case A - UV is treated as noise. Another issue is that SVD gives a factorization with negative entries. Non-negative matrix Factorization (NMF) is more appropriate in some contexts where we want to keep entries non-negative. NMF is discussed in Chapter 8.13

In the above setting, A was available fully and we wished to find U and V to identify the basic factors. However, in a case such as movie recommendations, each customer may have seen only a small fraction of the movies, so it may be more natural to assume that we are given just a few elements of A and wish to estimate A. If A was an arbitrary matrix of size $n \times d$, this would require $\Omega(nd)$ pieces of information and cannot be done with a few entries. But again hypothesize that A was a small rank matrix with added noise. If now we also assume that the given entries are randomly drawn according to some known distribution, then there is a possibility that SVD can be used to estimate the whole of A. This area is called collaborative filtering and one of its uses is to recommend movies or to target an ad to a customer based on one or two purchases. We do not describe it here.

3.9.3 Clustering a Mixture of Spherical Gaussians

Clustering is the task of partitioning a set of points into k subsets or clusters where each cluster consists of "nearby" points. Different definitions of the quality of a clustering lead to different solutions. Clustering is an important area which we will study in detail in Chapter ??. Here we will see how to solve a particular clustering problem using singular value decomposition.

Mathematical formulations of clustering tend to have the property that finding the highest quality solution to a given set of data is NP-hard. One way around this is to assume stochastic models of input data and devise algorithms to cluster data generated by such models. Mixture models are a very important class of stochastic models. A mixture is a probability density or distribution that is the weighted sum of simple component probability densities. It is of the form

$$F = w_1 p_1 + w_2 p_2 + \dots + w_k p_k$$

where p_1, p_2, \ldots, p_k are the basic probability densities and w_1, w_2, \ldots, w_k are positive real numbers called mixture weights that add up to one. Clearly, F is a probability density and integrates to one.

The model fitting problem is to fit a mixture of k basic densities to n independent, identically distributed samples, each sample drawn according to the same mixture distribution F. The class of basic densities is known, but various parameters such as their means and the component weights of the mixture are not. Here, we deal with the case where the basic densities are all spherical Gaussians. There are two equivalent ways of thinking of the sample generation process (which is hidden; only the samples are given):

- 1. Pick each sample according to the density F on \mathbf{R}^d .
- 2. Pick a random *i* from $\{1, 2, ..., k\}$ where probability of picking *i* is w_i . Then, pick a sample according to the density p_i .

One approach to the model-fitting problem is to break it into two subproblems:

- 1. First, cluster the set of samples into k clusters C_1, C_2, \ldots, C_k , where, C_i is the set of samples generated according to p_i (see (2) above) by the hidden generation process.
- 2. Then, fit a single Gaussian distribution to each cluster of sample points.

The second problem is relatively easier and indeed we saw the solution in Chapter (2), where we showed that taking the empirical mean (the mean of the sample) and the empirical standard deviation gives us the best-fit Gaussian. The first problem is harder and this is what we discuss here.

If the component Gaussians in the mixture have their centers very close together, then the clustering problem is unresolvable. In the limiting case where a pair of component densities are the same, there is no way to distinguish between them. What condition on the inter-center separation will guarantee unambiguous clustering? First, by looking at 1-dimensional examples, it is clear that this separation should be measured in units of the standard deviation, since the density is a function of the number of standard deviation from the mean. In one dimension, if two Gaussians have inter-center separation at least six times the maximum of their standard deviations, then they hardly overlap. This is summarized in the question: How many standard deviations apart are the means? In one dimension, if the answer is at least six, we can easily tell the Gaussians apart. What is the analog of this in higher dimensions? We discussed in Chapter (2) distances between two sample points from the same Gaussian as well the distance between two sample points from two different Gaussians. Recall from that discussion that if

• If x and y are two independent samples from the same spherical Gaussian with standard deviation¹¹ σ then

$$|\mathbf{x} - \mathbf{y}|^2 \approx 2(\sqrt{d} \pm O(1))^2 \sigma^2$$

• If **x** and **y** are samples from different spherical Gaussians each of standard deviation σ and means separated by distance Δ , then

$$|\mathbf{x} - \mathbf{y}|^2 \approx 2(\sqrt{d} \pm O(1))^2 \sigma^2 + \Delta^2.$$

To ensure that points from the same Gaussian are closer to each other than points from different Gaussians, we need

$$2(\sqrt{d} - O(1))^2 \sigma^2 + \Delta^2 > 2(\sqrt{d} + O(1))^2 \sigma^2.$$

Expanding the squares, the high order term 2d cancels and we need that

$$\Delta > cd^{1/4},$$

for some constant c. While this was not a completely rigorous argument, it can be used to show that a distance based clustering approach (see Chapter 2 for an example) requires an inter-mean separation of at least $cd^{1/4}$ standard deviations to succeed, thus unfortunately not keeping with mnemonic of a constant number of standard deviations separation of the means. Here, indeed, we will show that $\Omega(1)$ standard deviations suffice provided the number k of Gaussians is O(1).

The central idea is the following. Suppose we can find the subspace spanned by the k centers and project the sample points to this subspace. The projection of a spherical Gaussian with standard deviation σ remains a spherical Gaussian with standard deviation σ (Lemma 3.15). In the projection, the inter-center separation remains the same. So in the projection, the Gaussians are distinct provided the inter-center separation in the whole space is at least $ck^{1/4}\sigma$ which is a lot less than $cd^{1/4}\sigma$ for $k \ll d$. Interestingly, we will see that the subspace spanned by the k-centers is essentially the best-fit k-dimensional subspace that can be found by singular value decomposition.

Lemma 3.15 Suppose p is a d-dimensional spherical Gaussian with center μ and standard deviation σ . The density of p projected onto a k-dimensional subspace V is a spherical Gaussian with the same standard deviation.

¹¹Since a spherical Gaussian has the same standard deviation in every direction, we call it the standard deviation of the Gaussian.

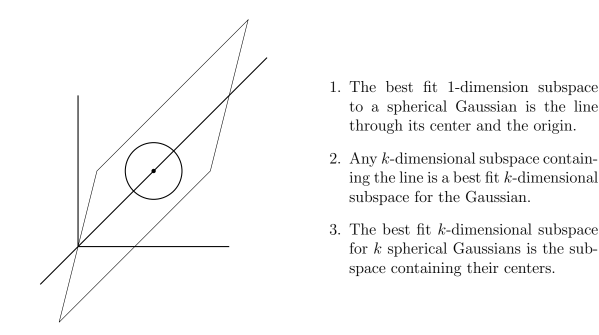


Figure 3.4: Best fit subspace to a spherical Gaussian.

Proof: Rotate the coordinate system so V is spanned by the first k coordinate vectors. The Gaussian remains spherical with standard deviation σ although the coordinates of its center have changed. For a point $\mathbf{x} = (x_1, x_2, \ldots, x_d)$, we will use the notation $\mathbf{x}' = (x_1, x_2, \ldots, x_k)$ and $\mathbf{x}'' = (x_{k+1}, x_{k+2}, \ldots, x_n)$. The density of the projected Gaussian at the point (x_1, x_2, \ldots, x_k) is

$$ce^{-\frac{|\mathbf{x}'-\boldsymbol{\mu}'|^2}{2\sigma^2}}\int\limits_{\mathbf{x}''}e^{-\frac{|\mathbf{x}''-\boldsymbol{\mu}''|^2}{2\sigma^2}}d\mathbf{x}''=c'e^{-\frac{|\mathbf{x}'-\boldsymbol{\mu}'|^2}{2\sigma^2}}.$$

This implies the lemma.

We now show that the top k singular vectors produced by the SVD span the space of the k centers. First, we extend the notion of best fit to probability distributions. Then we show that for a single spherical Gaussian whose center is not the origin, the best fit 1-dimensional subspace is the line though the center of the Gaussian and the origin. Next, we show that the best fit k-dimensional subspace for a single Gaussian whose center is not the origin is any k-dimensional subspace containing the line through the Gaussian's center and the origin. Finally, for k spherical Gaussians, the best fit k-dimensional subspace is the subspace containing their centers. Thus, the SVD finds the subspace that contains the centers.

Recall that for a set of points, the best-fit line is the line passing through the origin that maximizes the sum of squared lengths of the projections of the points onto the line. We extend this definition to probability densities instead of a set of points.

Definition 3.1 If p is a probability density in d space, the best fit line for p is the line $l = \{c\mathbf{v_1} : c \in R\}$ where

$$\mathbf{v_1} = \arg \max_{|\mathbf{v}|=1} E_{\mathbf{x} \sim p} \left[(\mathbf{v}^T \mathbf{x})^2 \right]$$

For a spherical Gaussian centered at the origin, it is easy to see that any line passing through the origin is a best fit line. Our next lemma shows that the best fit line for a spherical Gaussian centered at $\mu \neq 0$ is the line passing through μ and the origin.

Lemma 3.16 Let the probability density p be a spherical Gaussian with center $\mu \neq 0$. The unique best fit 1-dimensional subspace is the line passing through μ and the origin. If $\mu = 0$, then any line through the origin is a best-fit line.

Proof: For a randomly chosen \boldsymbol{x} (according to p) and a fixed unit length vector \mathbf{v} ,

$$\begin{split} E_{\mathbf{x}\sim p} \left[(\mathbf{v}^T \mathbf{x})^2 \right] &= E_{\mathbf{x}\sim p} \left[\left(\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) + \mathbf{v}^T \boldsymbol{\mu} \right)^2 \right] \\ &= E_{\mathbf{x}\sim p} \left[\left(\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) \right)^2 + 2 \left(\mathbf{v}^T \boldsymbol{\mu} \right) \left(\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) \right) + \left(\mathbf{v}^T \boldsymbol{\mu} \right)^2 \right] \\ &= E_{\mathbf{x}\sim p} \left[\left(\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) \right)^2 \right] + 2 \left(\mathbf{v}^T \boldsymbol{\mu} \right) E \left[\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) \right] + \left(\mathbf{v}^T \boldsymbol{\mu} \right)^2 \\ &= E_{\mathbf{x}\sim p} \left[\left(\mathbf{v}^T \left(\mathbf{x} - \boldsymbol{\mu} \right) \right)^2 \right] + \left(\mathbf{v}^T \boldsymbol{\mu} \right)^2 \\ &= \sigma^2 + \left(\mathbf{v}^T \boldsymbol{\mu} \right)^2 \end{split}$$

where the fourth line follows from the fact that $E[\mathbf{v}^T(\mathbf{x} - \boldsymbol{\mu})] = 0$, and the fifth line follows from the fact that $E[(\mathbf{v}^T(\mathbf{x} - \boldsymbol{\mu}))^2]$ is the variance in the direction \mathbf{v} . The best fit line \mathbf{v} maximizes $E_{\mathbf{x}\sim p}[(\mathbf{v}^T\mathbf{x})^2]$ and therefore maximizes $(\mathbf{v}^T\boldsymbol{\mu})^2$. This is maximized when \mathbf{v} is aligned with the center $\boldsymbol{\mu}$. To see uniqueness, just note that if $\boldsymbol{\mu} \neq 0$, then $\mathbf{v}^T\boldsymbol{\mu}$ is strictly less when \mathbf{v} is not aligned with the center.

We now extend Definition 3.1 to k-dimensional subspaces.

Definition 3.2 If p is a probability density in d-space then the best-fit k-dimensional subspace V_k is

$$V_k = \underset{V:dim(V)=k}{\operatorname{argmax}} \underset{\mathbf{x} \sim p}{E} \left[|\operatorname{proj}(\mathbf{x}, V)|^2 \right],$$

where $proj(\mathbf{x}, V)$ is the orthogonal projection of \mathbf{x} onto V.

Lemma 3.17 For a spherical Gaussian with center μ , a k-dimensional subspace is a best fit subspace if and only if it contains μ .

Proof: If $\mu = 0$, then by symmetry any k-dimensional subspace is a best-fit subspace. If $\mu \neq 0$, then, the best-fit line must pass through μ by Lemma (3.16). Now, as in the greedy algorithm for finding subsequent singular vectors, we would project perpendicular to the first singular vector. But after the projection, the mean of the Gaussian becomes 0 and any vectors will do as subsequent best-fit directions. This leads to the following theorem.

Theorem 3.18 If p is a mixture of k spherical Gaussians, then the best fit k-dimensional subspace contains the centers. In particular, if the means of the Gaussians are linearly independent, the space spanned by them is the unique best-fit k dimensional subspace.

Proof: Let p be the mixture $w_1p_1+w_2p_2+\cdots+w_kp_k$. Let V be any subspace of dimension k or less. Then,

$$\mathop{E}_{\mathbf{x} \sim p} \left[|\operatorname{proj}(\mathbf{x}, V)|^2 \right] = \sum_{i=1}^k w_i \mathop{E}_{\mathbf{x} \sim p_i} \left[|\operatorname{proj}(\mathbf{x}, V)|^2 \right]$$

If V contains the centers of the densities p_i , by Lemma 3.17, each term in the summation is individually maximized, which implies the entire summation is maximized, proving the theorem.

For an infinite set of points drawn according to the mixture, the k-dimensional SVD subspace gives exactly the space of the centers. In reality, we have only a large number of samples drawn according to the mixture. However, it is intuitively clear that as the number of samples increases, the set of sample points will approximate the probability density and so the SVD subspace of the sample will be close to the space spanned by the centers. The details of how close it gets as a function of the number of samples are technical and we do not carry this out here.

3.9.4 Ranking Documents and Web Pages

An important task for a document collection is to rank the documents according to their intrinsic relevance to the collection. A good candidate definition of "intrinsic relevance" is a document's projection onto the best-fit direction for that collection, namely the top left-singular vector of the term-document matrix. An intuitive reason for this is that this direction has the maximum sum of squared projections of the collection and so can be thought of as a synthetic term-document vector best representing the document collection.

Ranking in order of the projection of each document's term vector along the best fit direction has a nice interpretation in terms of the power method. For this, we consider a different example, that of the web with hypertext links. The World Wide Web can be represented by a directed graph whose nodes correspond to web pages and directed edges to hypertext links between pages. Some web pages, called *authorities*, are the most prominent sources for information on a given topic. Other pages called *hubs*, are ones that identify the authorities on a topic. Authority pages are pointed to by many hub pages and hub pages point to many authorities. One is led to what seems like a circular definition: a hub is a page that points to many authorities and an authority is a page that is pointed to by many hubs. One would like to assign hub weights and authority weights to each node of the web. If there are n nodes, the hub weights form an n-dimensional vector \mathbf{u} and the authority weights form an n-dimensional vector \mathbf{v} . Suppose A is the adjacency matrix representing the directed graph. Here a_{ij} is 1 if there is a hypertext link from page i to page j and 0 otherwise. Given hub vector \mathbf{u} , the authority vector \mathbf{v} could be computed by the formula

$$v_j \propto \sum_{i=1}^a u_i a_{ij}$$

since the right hand side is the sum of the hub weights of all the nodes that point to node j. In matrix terms,

$$\mathbf{v} = A^T \mathbf{u} / |A^T \mathbf{u}|.$$

Similarly, given an authority vector \mathbf{v} , the hub vector \mathbf{u} could be computed by $\mathbf{u} = A\mathbf{v}/|A\mathbf{v}|$. Of course, at the start, we have neither vector. But the above discussion suggests a power iteration. Start with any \mathbf{v} . Set $\mathbf{u} = A\mathbf{v}$, then set $\mathbf{v} = A^T\mathbf{u}$, then renormalize and repeat the process. We know from the power method that this converges to the left and right-singular vectors. So after sufficiently many iterations, we may use the left vector \mathbf{u} as the hub weights vector and project each column of A onto this direction and rank columns (authorities) in order of this projection. But the projections just form the vector $A^T\mathbf{u}$ which equals a multiple of \mathbf{v} . So we can just rank by order of the v_j . This is the basis of an algorithm called the HITS algorithm, which was one of the early proposals for ranking web pages.

A different ranking called *pagerank* is widely used. It is based on a random walk on the graph described above. We will study random walks in detail in Chapter 5.

3.9.5 An Application of SVD to a Discrete Optimization Problem

In clustering a mixture of Gaussians, SVD was used as a dimension reduction technique. It found a k-dimensional subspace (the space of centers) of a d-dimensional space and made the Gaussian clustering problem easier by projecting the data to the subspace. Here, instead of fitting a model to data, we consider an optimization problem where applying dimension reduction makes the problem easier. The use of SVD to solve discrete optimization problems is a relatively new subject with many applications. We start with an important NP-hard problem, the maximum cut problem for a directed graph G(V, E).

The maximum cut problem is to partition the nodes of an *n*-node directed graph into two subsets S and \overline{S} so that the number of edges from S to \overline{S} is maximized. Let A be the adjacency matrix of the graph. With each vertex i, associate an indicator variable x_i . The variable x_i will be set to 1 for $i \in S$ and 0 for $i \in \overline{S}$. The vector $\mathbf{x} = (x_1, x_2, \ldots, x_n)$ is unknown and we are trying to find it or equivalently the cut, so as to maximize the number of edges across the cut. The number of edges across the cut is precisely

$$\sum_{i,j} x_i (1 - x_j) a_{ij}$$

Thus, the maximum cut problem can be posed as the optimization problem

Maximize
$$\sum_{i,j} x_i(1-x_j)a_{ij}$$
 subject to $x_i \in \{0,1\}$.

In matrix notation,

$$\sum_{i,j} x_i (1 - x_j) a_{ij} = \mathbf{x}^T A (\mathbf{1} - \mathbf{x}),$$

where **1** denotes the vector of all 1's. So, the problem can be restated as

Maximize
$$\mathbf{x}^T A(\mathbf{1} - \mathbf{x})$$
 subject to $x_i \in \{0, 1\}.$ (3.1)

This problem is NP-hard. However we will see that for dense graphs, that is, graphs with $\Omega(n^2)$ edges and therefore whose optimal solution has size $\Omega(n^2)$,¹² we can use the SVD to find a near optimal solution in polynomial time. To do so we will begin by computing the SVD of A and replacing A by $A_k = \sum_{i=1}^k \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ in (3.1) to get

Maximize
$$\mathbf{x}^T A_k(\mathbf{1} - \mathbf{x})$$
 subject to $x_i \in \{0, 1\}.$ (3.2)

Note that the matrix A_k is no longer a 0-1 adjacency matrix.

We will show that:

- 1. For each 0-1 vector \mathbf{x} , $\mathbf{x}^T A_k(\mathbf{1} \mathbf{x})$ and $\mathbf{x}^T A(\mathbf{1} \mathbf{x})$ differ by at most $\frac{n^2}{\sqrt{k+1}}$. Thus, the maxima in (3.1) and (3.2) differ by at most this amount.
- 2. A near optimal **x** for (3.2) can be found in time $n^{O(k)}$ by exploiting the low rank of A_k , which is polynomial time for constant k. By Item 1 this is near optimal for (3.1) where near optimal means with additive error of at most $\frac{n^2}{\sqrt{k+1}}$.

First, we prove Item 1. Since \mathbf{x} and $\mathbf{1} - \mathbf{x}$ are 0-1 *n*-vectors, each has length at most \sqrt{n} . By the definition of the 2-norm, $|(A - A_k)(\mathbf{1} - \mathbf{x})| \leq \sqrt{n}||A - A_k||_2$. Now since $\mathbf{x}^T (A - A_k)(\mathbf{1} - \mathbf{x})$ is the dot product of the vector \mathbf{x} with the vector $(A - A_k)(\mathbf{1} - \mathbf{x})$,

$$|\mathbf{x}^T (A - A_k)(\mathbf{1} - \mathbf{x})| \le n ||A - A_k||_2.$$

By Lemma 3.8, $||A - A_k||_2 = \sigma_{k+1}(A)$. The inequalities,

$$(k+1)\sigma_{k+1}^2 \le \sigma_1^2 + \sigma_2^2 + \cdots + \sigma_{k+1}^2 \le ||A||_F^2 = \sum_{i,j} a_{ij}^2 \le n^2$$

imply that $\sigma_{k+1}^2 \leq \frac{n^2}{k+1}$ and hence $||A - A_k||_2 \leq \frac{n}{\sqrt{k+1}}$ proving Item 1.

¹²Any graph of m edges has a cut of size at least m/2. This can be seen by noting that the expected size of the cut for a random $\mathbf{x} \in \{0, 1\}^n$ is exactly m/2.

Next we focus on Item 2. It is instructive to look at the special case when k=1 and A is approximated by the rank one matrix A_1 . An even more special case when the left and right-singular vectors **u** and **v** are identical is already NP-hard to solve exactly because it subsumes the problem of whether for a set of n integers, $\{a_1, a_2, \ldots, a_n\}$, there is a partition into two subsets whose sums are equal. However, for that problem, there is an efficient dynamic programming algorithm that finds a near-optimal solution. We will build on that idea for the general rank k problem.

For Item 2, we want to maximize $\sum_{i=1}^{k} \sigma_i(\mathbf{x}^T \mathbf{u}_i)(\mathbf{v}_i^T(\mathbf{1} - \mathbf{x}))$ over 0-1 vectors \mathbf{x} . A piece of notation will be useful. For any $S \subseteq \{1, 2, ..., n\}$, write $\mathbf{u}_i(S)$ for the sum of coordinates of the vector \mathbf{u}_i corresponding to elements in the set S, that is, $\mathbf{u}_i(S) = \sum_{j \in S} u_{ij}$, and similarly for \mathbf{v}_i . We will find S to maximize $\sum_{i=1}^{k} \sigma_i \mathbf{u}_i(S) \mathbf{v}_i(\bar{S})$ using dynamic programming.

For a subset S of $\{1, 2, ..., n\}$, define the 2k-dimensional vector

$$\mathbf{w}(S) = (\mathbf{u}_1(S), \mathbf{v}_1(\bar{S}), \mathbf{u}_2(S), \mathbf{v}_2(\bar{S}), \dots, \mathbf{u}_k(S), \mathbf{v}_k(\bar{S}))$$

If we had the list of all such vectors, we could find $\sum_{i=1}^{k} \sigma_i \mathbf{u}_i(S) \mathbf{v}_i(\bar{S})$ for each of them and take the maximum. There are 2^n subsets S, but several S could have the same $\mathbf{w}(S)$ and in that case it suffices to list just one of them. Round each coordinate of each \mathbf{u}_i to the nearest integer multiple of $\frac{1}{nk^2}$. Call the rounded vector $\tilde{\mathbf{u}}_i$. Similarly obtain $\tilde{\mathbf{v}}_i$. Let $\tilde{\mathbf{w}}(S)$ denote the vector $(\tilde{\mathbf{u}}_1(S), \tilde{\mathbf{v}}_1(\bar{S}), \tilde{\mathbf{u}}_2(S), \tilde{\mathbf{v}}_2(\bar{S}), \dots, \tilde{\mathbf{u}}_k(S), \tilde{\mathbf{v}}_k(\bar{S}))$. We will construct a list of all possible values of the vector $\tilde{\mathbf{w}}(S)$. Again, if several different S's lead to the same vector $\tilde{\mathbf{w}}(S)$, we will keep only one copy on the list. The list will be constructed by dynamic programming. For the recursive step, assume we already have a list of all such vectors for $S \subseteq \{1, 2, \dots, i\}$ and wish to construct the list for $S \subseteq \{1, 2, \dots, i+1\}$. Each $S \subseteq \{1, 2, \dots, i\}$ leads to two possible $S' \subseteq \{1, 2, \dots, i+1\}$, namely, S and $S \cup \{i+1\}$. In the first case, the vector $\tilde{\mathbf{w}}(S') = (\tilde{\mathbf{u}}_1(S), \tilde{\mathbf{v}}_1(\bar{S}) + \tilde{v}_{1,i+1}, \tilde{\mathbf{u}}_2(S), \tilde{\mathbf{v}}_2(\bar{S}) + \tilde{v}_{2,i+1}, \dots, \dots)$. In the second case, it is $\tilde{\mathbf{w}}(S') = (\tilde{\mathbf{u}}_1(S) + \tilde{u}_{1,i+1}, \tilde{\mathbf{v}}_1(\bar{S}), \tilde{\mathbf{u}}_2(S) + \tilde{u}_{2,i+1}, \tilde{\mathbf{v}}_2(\bar{S}), \dots, \dots)$. We put in these two vectors for each vector in the previous list. Then, crucially, we prune -i.e., eliminate duplicates.

Assume that k is constant. Now, we show that the error is at most $\frac{n^2}{\sqrt{k+1}}$ as claimed. Since $\mathbf{u_i}$ and $\mathbf{v_i}$ are unit length vectors, $|\mathbf{u_i}(S)|, |\mathbf{v_i}(\bar{S})| \leq \sqrt{n}$. Also $|\mathbf{\tilde{u_i}}(S) - \mathbf{u_i}(S)| \leq \frac{n}{nk^2} = \frac{1}{k^2}$ and similarly for $\mathbf{v_i}$. To bound the error, we use an elementary fact: if a and b are reals with $|a|, |b| \leq M$ and we estimate a by a' and b by b' so that $|a - a'|, |b - b'| \leq \delta \leq M$, then a'b' is an estimate of ab in the sense

$$|ab - a'b'| = |a(b - b') + b'(a - a')| \le |a||b - b'| + (|b| + |b - b'|)|a - a'| \le 3M\delta.$$

Using this, we get that

$$\left|\sum_{i=1}^{k} \sigma_i \tilde{\mathbf{u}}_i(S) \tilde{\mathbf{v}}_i(\bar{S}) - \sum_{i=1}^{k} \sigma_i \mathbf{u}_i(S) \mathbf{v}_i(S)\right| \le 3k\sigma_1 \sqrt{n}/k^2 \le 3n^{3/2}/k \le n^2/k,$$

and this meets the claimed error bound.

Next, we show that the running time is polynomially bounded. First, $|\tilde{\mathbf{u}}_{\mathbf{i}}(S)|, |\tilde{\mathbf{v}}_{\mathbf{i}}(S)| \leq 2\sqrt{n}$. Since $\tilde{\mathbf{u}}_{\mathbf{i}}(S)$ and $\tilde{\mathbf{v}}_{\mathbf{i}}(S)$ are all integer multiples of $1/(nk^2)$, there are at most $2n^{3/2}k^2$ possible values of $\tilde{\mathbf{u}}_{\mathbf{i}}(S)$ and $\tilde{\mathbf{v}}_{\mathbf{i}}(S)$ from which it follows that the list of $\tilde{\mathbf{w}}(S)$ never gets larger than $(2n^{3/2}k^2)^{2k}$ which for fixed k is polynomially bounded.

We summarize what we have accomplished.

Theorem 3.19 Given a directed graph G(V, E), a cut of size at least the maximum cut minus $O\left(\frac{n^2}{\sqrt{k}}\right)$ can be computed in time polynomial in n for any fixed k.

Note that achieving the same accuracy in time polynomial in n and k would give an exact max cut in polynomial time.

3.10 Bibliographic Notes

Singular value decomposition is fundamental to numerical analysis and linear algebra. There are many texts on these subjects and the interested reader may want to study these. A good reference is [GvL96]. The material on clustering a mixture of Gaussians in Section 3.9.3 is from [VW02]. Modeling data with a mixture of Gaussians is a standard tool in statistics. Several well-known heuristics like the expectation-minimization algorithm are used to learn (fit) the mixture model to data. Recently, in theoretical computer science, there has been modest progress on provable polynomial-time algorithms for learning mixtures. Some references are [DS07], [AK], [AM05], and [MV10]. The application to the discrete optimization problem is from [FK99]. The section on ranking documents/webpages is from two influential papers, one on hubs and authorities by Jon Kleinberg [Kle99] and the other on pagerank by Page, Brin, Motwani and Winograd [BMPW98].

3.11 Exercises

Exercise 3.1 (Least squares vertical error) In many experiments one collects the value of a parameter at various instances of time. Let y_i be the value of the parameter y at time x_i . Suppose we wish to construct the best linear approximation to the data in the sense that we wish to minimize the mean square error. Here error is measured vertically rather than perpendicular to the line. Develop formulas for m and b to minimize the mean square error of the points $\{(x_i, y_i) | 1 \le i \le n\}$ to the line y = mx + b.

Exercise 3.2 Given five observed variables, height, weight, age, income, and blood pressure of n people, how would one find the best least squares fit affine subspace of the form

 a_1 (height) + a_2 (weight) + a_3 (age) + a_4 (income) + a_5 (blood pressure) = a_6

Here a_1, a_2, \ldots, a_6 are the unknown parameters. If there is a good best fit 4-dimensional affine subspace, then one can think of the points as lying close to a 4-dimensional sheet rather than points lying in 5-dimensions. Why might it be better to use the perpendicular distance to the affine subspace rather than vertical distance where vertical distance is measured along the coordinate axis corresponding to one of the variables?

Exercise 3.3 Manually find the best fit lines (not subspaces which must contain the origin) through the points in the sets below. Subtract the center of gravity of the points in the set from each of the points in the set and find the best fit line for the resulting points. Does the best fit line for the original data go through the origin?

- 1. (4,4) (6,2)
- 2. (4,2) (4,4) (6,2) (6,4)
- 3. (3,2.5) (3,5) (5,1) (5,3.5)

Exercise 3.4 Manually determine the best fit line through the origin for each of the following sets of points. Is the best fit line unique? Justify your answer for each of the subproblems.

- 1. $\{(0,1),(1,0)\}$
- 2. $\{(0,1),(2,0)\}$

Exercise 3.5 Manually find the left and right-singular vectors, the singular values, and the SVD decomposition of the matrices in Figure 3.5.

Exercise 3.6 Consider the matrix

$$A = \begin{pmatrix} 1 & 2\\ -1 & 2\\ 1 & -2\\ -1 & -2 \end{pmatrix}$$

$$(0,3) \bullet M = \begin{pmatrix} 1 & 1 \\ 0 & 3 \\ 3 & 0 \end{pmatrix} (0,2) \bullet (1,3) \bullet (1,3)$$

Figure 3.5: SVD problem

- 1. Run the power method starting from $x = \begin{pmatrix} 1 \\ 1 \end{pmatrix}$ for k = 3 steps. What does this give as an estimate of v_1 ?
- 2. What actually are the v_i 's, σ_i 's, and u_i 's? It may be easiest to do this by computing the eigenvectors of $B = A^T A$.
- 3. Suppose matrix A is a database of restaurant ratings: each row is a person, each column is a restaurant, and a_{ij} represents how much person i likes restaurant j. What might v_1 represent? What about u_1 ? How about the gap $\sigma_1 \sigma_2$?

Exercise 3.7 Let A be a square $n \times n$ matrix whose rows are orthonormal. Prove that the columns of A are orthonormal.

Exercise 3.8 Suppose A is a $n \times n$ matrix with block diagonal structure with k equal size blocks where all entries of the *i*th block are a_i with $a_1 > a_2 > \cdots > a_k > 0$. Show that A has exactly k nonzero singular vectors $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_k}$ where $\mathbf{v_i}$ has the value $(\frac{k}{n})^{1/2}$ in the coordinates corresponding to the *i*th block and 0 elsewhere. In other words, the singular vectors exactly identify the blocks of the diagonal. What happens if $a_1 = a_2 = \cdots = a_k$? In the case where the a_i are equal, what is the structure of the set of all possible singular vectors?

Hint: By symmetry, the top singular vector's components must be constant in each block.

Exercise 3.9 Interpret the first right and left-singular vectors for the document term matrix.

Exercise 3.10 Verify that the sum of r-rank one matrices $\sum_{i=1}^{r} c_i \mathbf{x}_i \mathbf{y}_i^T$ can be written as XCY^T , where the \mathbf{x}_i are the columns of X, the \mathbf{y}_i are the columns of Y, and C is a diagonal matrix with the constants c_i on the diagonal.

Exercise 3.11 Let $\sum_{i=1}^{r} \sigma_i \mathbf{u}_i \mathbf{v}_i^T$ be the SVD of A. Show that $|\mathbf{u}_1^T A| = \max_{|\mathbf{u}|=1} |\mathbf{u}^T A|$ equals σ_1 .

Exercise 3.12 If $\sigma_1, \sigma_2, \ldots, \sigma_r$ are the singular values of A and $\mathbf{v}_1, \mathbf{v}_2, \ldots, \mathbf{v}_r$ are the corresponding right-singular vectors, show that

- 1. $A^T A = \sum_{i=1}^r \sigma_i^2 \mathbf{v_i v_i}^T$
- 2. $\mathbf{v}_1, \mathbf{v}_2, \dots \mathbf{v}_r$ are eigenvectors of $A^T A$.
- 3. Assuming that the eigenvectors of $A^T A$ are unique up to multiplicative constants, conclude that the singular vectors of A (which by definition must be unit length) are unique up to sign.

Exercise 3.13 Let $\sum_{i} \sigma_{i} u_{i} v_{i}^{T}$ be the singular value decomposition of a rank r matrix A. Let $A_{k} = \sum_{i=1}^{k} \sigma_{i} u_{i} v_{i}^{T}$ be a rank k approximation to A for some k < r. Express the following quantities in terms of the singular values $\{\sigma_{i}, 1 \leq i \leq r\}$.

- 1. $||A_k||_F^2$
- 2. $||A_k||_2^2$
- 3. $||A A_k||_F^2$
- 4. $||A A_k||_2^2$

Exercise 3.14 If A is a symmetric matrix with distinct singular values, show that the left and right singular vectors are the same and that $A = VDV^T$.

Exercise 3.15 Let A be a matrix. How would you compute

$$\mathbf{v_1} = \operatorname*{arg\,max}_{|\mathbf{v}|=1} |A\mathbf{v}|?$$

How would you use or modify your algorithm for finding $\mathbf{v_1}$ to compute the first few singular vectors of A.

Exercise 3.16 Use the power method to compute the singular value decomposition of the matrix

$$A = \left(\begin{array}{cc} 1 & 2\\ 3 & 4 \end{array}\right)$$

Exercise 3.17 Write a program to implement the power method for computing the first singular vector of a matrix. Apply your program to the matrix

$$A = \begin{pmatrix} 1 & 2 & 3 & \cdots & 9 & 10 \\ 2 & 3 & 4 & \cdots & 10 & 0 \\ \vdots & \vdots & \vdots & & & \vdots \\ 9 & 10 & 0 & \cdots & 0 & 0 \\ 10 & 0 & 0 & \cdots & 0 & 0 \end{pmatrix}.$$

Exercise 3.18 Modify the power method to find the first four singular vectors of a matrix A as follows. Randomly select four vectors and find an orthonormal basis for the space spanned by the four vectors. Then multiple each of the basis vectors times A and find a new orthonormal basis for the space spanned by the resulting four vectors. Apply your method to find the first four singular vectors of matrix A of Exercise 3.17. In Matlab the command orth finds an orthonormal basis for the space spanned by a set of vectors.

Exercise 3.19 A matrix M is positive semi-definite if for all $\mathbf{x}, \mathbf{x}^T M \mathbf{x} \ge 0$.

- 1. Let A be a real valued matrix. Prove that $B = AA^T$ is positive semi-definite.
- 2. Let A be the adjacency matrix of a graph. The Laplacian of A is L = D A where D is a diagonal matrix whose diagonal entries are the row sums of A. Prove that L is positive semi definite by showing that $L = B^T B$ where B is an m-by-n matrix with a row for each edge in the graph, a column for each vertex, and we define

$$b_{ei} = \begin{cases} -1 & \text{if } i \text{ is the endpoint of } e \text{ with lesser index} \\ 1 & \text{if } i \text{ is the endpoint of } e \text{ with greater index} \\ 0 & \text{if } i \text{ is not an endpoint of } e \end{cases}$$

Exercise 3.20 Prove that the eigenvalues of a symmetric real valued matrix are real.

Exercise 3.21 Suppose A is a square invertible matrix and the SVD of A is $A = \sum_{i} \sigma_{i} u_{i} v_{i}^{T}$. Prove that the inverse of A is $\sum_{i} \frac{1}{\sigma_{i}} v_{i} u_{i}^{T}$.

Exercise 3.22 Suppose A is square, but not necessarily invertible and has SVD $A = \sum_{i=1}^{r} \sigma_i u_i v_i^T$. Let $B = \sum_{i=1}^{r} \frac{1}{\sigma_i} v_i u_i^T$. Show that $BA\mathbf{x} = \mathbf{x}$ for all \mathbf{x} in the span of the right-singular vectors of A. For this reason B is sometimes called the pseudo inverse of A and can play the role of A^{-1} in many applications.

Exercise 3.23

- 1. For any matrix A, show that $\sigma_k \leq \frac{||A||_F}{\sqrt{k}}$.
- 2. Prove that there exists a matrix B of rank at most k such that $||A B||_2 \leq \frac{||A||_F}{\sqrt{k}}$.
- 3. Can the 2-norm on the left hand side in (b) be replaced by Frobenius norm?

Exercise 3.24 Suppose an $n \times d$ matrix A is given and you are allowed to preprocess A. Then you are given a number of d-dimensional vectors $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_m}$ and for each of these vectors you must find the vector $A\mathbf{x_j}$ approximately, in the sense that you must find a vector $\mathbf{y_j}$ satisfying $|\mathbf{y_j} - A\mathbf{x_j}| \leq \varepsilon ||A||_F |\mathbf{x_j}|$. Here $\varepsilon > 0$ is a given error bound. Describe an algorithm that accomplishes this in time $O\left(\frac{d+n}{\varepsilon^2}\right)$ per $\mathbf{x_j}$ not counting the preprocessing time. Hint: use Exercise 3.23.

Exercise 3.25 (Document-Term Matrices): Suppose we have an $m \times n$ document-term matrix A where each row corresponds to a document and has been normalized to length one. Define the "similarity" between two such documents by their dot product.

- 1. Consider a "synthetic" document whose sum of squared similarities with all documents in the matrix is as high as possible. What is this synthetic document and how would you find it?
- 2. How does the synthetic document in (1) differ from the center of gravity?
- 3. Building on (1), given a positive integer k, find a set of k synthetic documents such that the sum of squares of the mk similarities between each document in the matrix and each synthetic document is maximized. To avoid the trivial solution of selecting k copies of the document in (1), require the k synthetic documents to be orthogonal to each other. Relate these synthetic documents to singular vectors.
- 4. Suppose that the documents can be partitioned into k subsets (often called clusters), where documents in the same cluster are similar and documents in different clusters are not very similar. Consider the computational problem of isolating the clusters. This is a hard problem in general. But assume that the terms can also be partitioned into k clusters so that for i ≠ j, no term in the ith cluster occurs in a document in the jth cluster. If we knew the clusters and arranged the rows and columns in them to be contiguous, then the matrix would be a block-diagonal matrix. Of course the clusters are not known. By a "block" of the document-term matrix, we mean a submatrix with rows corresponding to the ith cluster of documents and columns corresponding to the ith cluster of terms. We can also partition any n vector into blocks. Show that any right-singular vector of the matrix must have the property that each of its blocks is a right-singular vector of the corresponding block of the document-term matrix.
- 5. Suppose now that the singular values of all the blocks are distinct (also across blocks). Show how to solve the clustering problem.

Hint: (4) Use the fact that the right-singular vectors must be eigenvectors of $A^T A$. Show that $A^T A$ is also block-diagonal and use properties of eigenvectors.

Exercise 3.26 Show that maximizing $\mathbf{x}^T \mathbf{u} \mathbf{u}^T (\mathbf{1} - \mathbf{x})$ subject to $x_i \in \{0, 1\}$ is equivalent to partitioning the coordinates of \mathbf{u} into two subsets where the sum of the elements in both subsets are as equal as possible.

Exercise 3.27 Read in a photo and convert to a matrix. Perform a singular value decomposition of the matrix. Reconstruct the photo using only 5%, 10%, 25%, 50% of the singular values.

1. Print the reconstructed photo. How good is the quality of the reconstructed photo?

2. What percent of the Forbenius norm is captured in each case?

Hint: If you use Matlab, the command to read a photo is imread. The types of files that can be read are given by imformats. To print the file use imwrite. Print using jpeg format. To access the file afterwards you may need to add the file extension .jpg. The command imread will read the file in uint8 and you will need to convert to double for the SVD code. Afterwards you will need to convert back to uint8 to write the file. If the photo is a color photo you will get three matrices for the three colors used.

- Exercise 3.28 1. Create a 100 × 100 matrix of random numbers between 0 and 1 such that each entry is highly correlated with the adjacency entries. Find the SVD of A. What fraction of the Frobenius norm of A is captured by the top 10 singular vectors? How many singular vectors are required to capture 95% of the Frobenius norm?
 - 2. Repeat (1) with a 100×100 matrix of statistically independent random numbers between 0 and 1.

Exercise 3.29 Show that the running time for the maximum cut algorithm in Section ?? can be carried out in time $O(n^3 + poly(n)k^k)$, where poly is some polynomial.

Exercise 3.30 Let $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_n}$ be *n* points in *d*-dimensional space and let *X* be the $n \times d$ matrix whose rows are the *n* points. Suppose we know only the matrix *D* of pairwise distances between points and not the coordinates of the points themselves. The set of points $\mathbf{x_1}, \mathbf{x_2}, \ldots, \mathbf{x_n}$ giving rise to the distance matrix *D* is not unique since any translation, rotation, or reflection of the coordinate system leaves the distances invariant. Fix the origin of the coordinate system so that the centroid of the set of points is at the origin. That is, $\sum_{i=1}^{n} \mathbf{x_i} = 0$.

1. Show that the elements of XX^T are given by

$$\mathbf{x}_{i}\mathbf{x}_{j}^{T} = -\frac{1}{2} \left[d_{ij}^{2} - \frac{1}{n} \sum_{j=1}^{n} d_{ij}^{2} - \frac{1}{n} \sum_{i=1}^{n} d_{ij}^{2} + \frac{1}{n^{2}} \sum_{i=1}^{n} \sum_{j=1}^{n} d_{ij}^{2} \right].$$

2. Describe an algorithm for determining the matrix X whose rows are the $\mathbf{x_i}$.

Exercise 3.31

1. Consider the pairwise distance matrix for twenty US cities given below. Use the algorithm of Exercise 3.30 to place the cities on a map of the US. The algorithm is called classical multidimensional scaling, cmdscale, in Matlab. Alternatively use the pairwise distance matrix to place the cities on a map of China.

Note: Any rotation or a mirror image of the map will have the same pairwise distances.

2. Suppose you had airline distances for 50 cities around the world. Could you use these distances to construct a 3-dimensional world model?

	В	В	С	D		о н	I	M	N	1	М
	ō	Ū		Ā						I	Ι
	Š	Ē				-		Μ			M
Boston	_	400		1551		9 1605	2596				
Buffalo	400	_									
Chicago	851	454	-	803	92	0 940			118		
Dallas	1551	1198									
Denver	1769	1370		663		- 879		879	172		
Houston	1605	1286	940	225	87			484	96	8 105	56
Los Angeles	2596	2198	1745	1240	83	1 1374	_	- 1603	233	9 152	24
Memphis	1137	803	482	420			1603	3 -	87	2 69	99
Miami	1255	1181	1188	1111	172	6 968	2339	872		- 151	1
Minneapolis	1123	731	355	862	70	0 1056	1524	699	151	1	-
New York	188	292	713	1374	163	1 1420	2451	957	109	2 101	18
Omaha	1282	883	432	586	48	8 794	1315	5 529	139	7 29	90
Philadelphia	271	279	666	1299	157	9 1341	2394	881	101	9 98	35
Phoenix	2300	1906	1453	887	58	6 1017	357	7 1263	198	2 128	30
Pittsburgh	483	178	410	1070	132	0 1137	2136	660	101	0 74	13
Saint Louis	1038	662	262	547	79	6 679	1589) 240	106	1 46	66
Salt Lake City	2099	1699	1260	999	37	1 1200	579) 1250	208	9 98	37
San Francisco	2699	2300	1858	1483	94	9 1645	347	7 1802	259	4 158	34
Seattle	2493	2117	1737	1681	102	1 1891	959) 1867	273	4 139	95
Washington D.C.	393	292	597	1185	149	4 1220	2300) 765	92	3 93	34
		Ν	О	Р	Р	Р	\mathbf{S}	\mathbf{S}	S	S	D
		Υ	Μ	Н	Η	Ι	\mathbf{t}	\mathbf{L}	\mathbf{F}	\mathbf{E}	\mathbf{C}
			А	Ι	Ο	Т	\mathbf{L}	\mathbf{C}		А	
Boston		188	1282	271	2300	483	1038	2099	2699	2493	393
Buffalo		292	883	279	1906	178	662	1699	2300	2117	292
Chicago		713	432	666	1453	410	262	1260	1858	1737	597
Dallas		1374	586	1299	887	1070	547	999	1483	1681	1185
Denver		1631	488	1579	586	1320	796	371	949	1021	1494
Houston		1420	794	1341	1017	1137	679	1200	1645	1891	1220
Los Angeles		2451		2394	357	2136	1589	579	347	959	2300
Memphis		957	529	881	1263	660	240	1250	1802	1867	765
Miami		1092		1019	1982	1010	1061	2089	2594	2734	923
Minneapolis		1018	290	985	1280	743	466	987	1584	1395	934
New York		-	1144	83	2145	317	875	1972	2571	2408	230
Omaha		1144		1094	1036	836	354	833	1429	1369	1014
Philadelphia		83	1094	-	2083	259	811	1925	2523	2380	123
Phoenix		2145		2083	-	1828	1272	504	653	1114	1973
Pittsburgh		317	836	259	1828	-	559	1668	2264	2138	192
Saint Louis		875	354	811	1272	559	-	1162	1744	1724	712
Salt Lake City		1972		1925	504	1668	1162	-	600	701	1848
San Francisco		2571		2523	653	2264	1744	600	-	678	2442
Seattle		2408		2380	1114	2138	1724	701	678	-	2329
Washington D	.C.	230	1014	123	1973	192	712	1848	2442	2329	-

City	Bei- jing	Tian- jin	Shang- hai	Chong- qing	Hoh- hot	Urum- qi	Lha- sa	Yin- chuan	Nan- ning	Har- bin	Chang- chun	Shen- yang
Beijing	0	125	1239	3026	480	3300	3736	1192	2373	1230	979	684
Tianjin	125	0	1150	1954	604	3330	3740	1316	2389	1207	955	661
Shanghai	1239	1150	0	1945	1717	3929	4157	2092	1892	2342	2090	1796
Chongqing	3026	1954	1945	0	1847	3202	2457	1570	993	3156	2905	2610
Hohhot	480	604	1717	1847	0	2825	3260	716	2657	1710	1458	1164
Urumqi	3300	3330	3929	3202	2825	0	2668	2111	4279	4531	4279	3985
Lhasa	3736	3740	4157	2457	3260	2668	0	2547	3431	4967	4715	4421
Yinchuan	1192	1316	2092	1570	716	2111	2547	0	2673	2422	2170	1876
Nanning	2373	2389	1892	993	2657	4279	3431	2673	0	3592	3340	3046
Harbin	1230	1207	2342	3156	1710	4531	4967	2422	3592	0	256	546
Changchun	979	955	2090	2905	1458	4279	4715	2170	3340	256	0	294
Shenyang	684	661	1796	2610	1164	3985	4421	1876	3046	546	294	0

4 Random Graphs

Large graphs appear in many contexts such as the World Wide Web, the internet, social networks, journal citations, and other places. What is different about the modern study of large graphs from traditional graph theory and graph algorithms is that here one seeks statistical properties of these very large graphs rather than an exact answer to questions. This is akin to the switch physics made in the late 19^{th} century in going from mechanics to statistical mechanics. Just as the physicists did, one formulates abstract models of graphs that are not completely realistic in every situation, but admit a nice mathematical development that can guide what happens in practical situations. Perhaps the most basic model is the G(n, p) model of a random graph. In this chapter, we study properties of the G(n, p) model as well as other models.

4.1 The G(n, p) Model

The G(n, p) model, due to Erdös and Rényi, has two parameters, n and p. Here n is the number of vertices of the graph and p is the edge probability. For each pair of distinct vertices, v and w, p is the probability that the edge (v,w) is present. The presence of each edge is statistically independent of all other edges. The graph-valued random variable with these parameters is denoted by G(n, p). When we refer to "the graph G(n, p)", we mean one realization of the random variable. In many cases, p will be a function of nsuch as p = d/n for some constant d. In this case, the expected degree of a vertex of the graph is $\frac{d}{n}(n-1) \approx d$. The interesting thing about the G(n, p) model is that even though edges are chosen independently with no "collusion", certain global properties of the graph emerge from the independent choices. For small p, with p = d/n, d < 1, each connected component in the graph is small. For d > 1, there is a giant component consisting of a constant fraction of the vertices. In addition, there is a rapid transition at the threshold d = 1. Below the threshold, the probability of a giant component is very small, and above the threshold, the probability is almost one.

The phase transition at the threshold d = 1 from very small o(n) size components to a giant $\Omega(n)$ sized component is illustrated by the following example. Suppose the vertices represent people and an edge means the two people it connects know each other. Given a

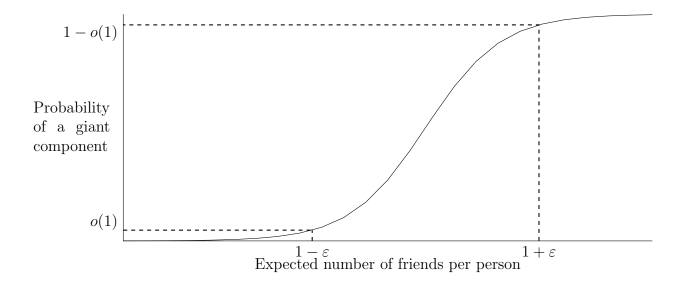


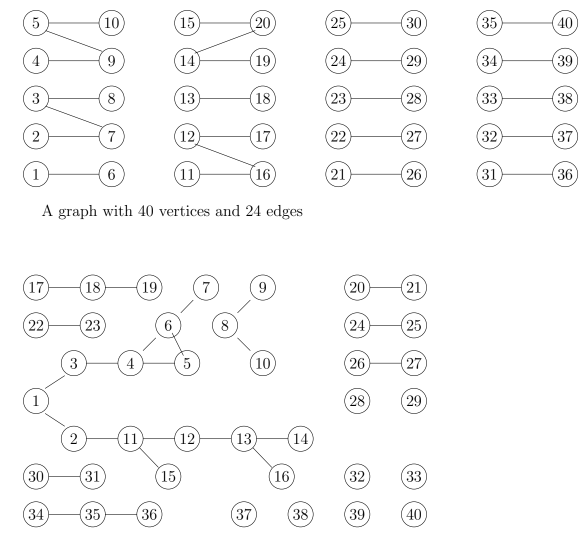
Figure 4.1: Probability of a giant component as a function of the expected number of people each person knows directly.

chain of connections, such as A knows B, B knows C, C knows D, ..., and Y knows Z, we say that A indirectly knows Z. Thus, all people belonging to a connected component of the graph indirectly know each other. Suppose each pair of people, independent of other pairs, tosses a coin that comes up heads with probability p = d/n. If it is heads, they know each other; if it comes up tails, they don't. The value of d can be interpreted as the expected number of people a single person directly knows. The question arises as to how large are sets of people who indirectly know each other ?

If the expected number of people each person knows is more than one, then a giant component of people, all of whom indirectly know each other, will be present consisting of a constant fraction of all the people. On the other hand, if in expectation, each person knows less than one person, the largest set of people who know each other indirectly is a vanishingly small fraction of the whole. Furthermore, the transition from the vanishing fraction to a constant fraction of the whole, happens abruptly between d slightly less than one to d slightly more than one. See Figure 4.1. Note that there is no global coordination of who knows whom. Each pair of individuals decides independently. Indeed, many large real-world graphs, with constant average degree, have a giant component. This is perhaps the most important global property of the G(n, p) model.

4.1.1 Degree Distribution

One of the simplest quantities to observe in a real graph is the number of vertices of given degree, called the vertex degree distribution. It is also very simple to study these distributions in G(n, p) since the degree of each vertex is the sum of n - 1 independent random variables, which results in a binomial distribution. Since we will be dealing with



A randomly generated G(n, p) graph with 40 vertices and 24 edges

Figure 4.2: Two graphs, each with 40 vertices and 24 edges. The second graph was randomly generated using the G(n,p) model with p = 1.2/n. A graph similar to the top graph is almost surely not going to be randomly generated in the G(n,p) model, whereas a graph similar to the lower graph will almost surely occur. Note that the lower graph consists of a giant component along with a number of small components that are trees.

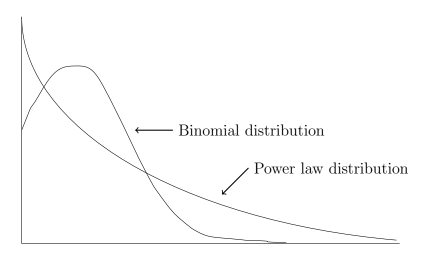


Figure 4.3: Illustration of the binomial and the power law distributions.

graphs where the number of vertices n, is large, from here on we often replace n-1 by n to simplify formulas.

Example: In $G(n, \frac{1}{2})$, each vertex is of degree close to n/2. In fact, for any $\varepsilon > 0$, the degree of each vertex almost surely is within $1 \pm \varepsilon$ times n/2. To see this, note that the probability that a vertex is of degree k is

$$\operatorname{Prob}\left(k\right) = \binom{n-1}{k} \left(\frac{1}{2}\right)^{k} \left(\frac{1}{2}\right)^{n-k} \approx \binom{n}{k} \left(\frac{1}{2}\right)^{k} \left(\frac{1}{2}\right)^{n-k} = \frac{1}{2^{n}} \binom{n}{k}.$$

This probability distribution has a mean m = n/2 and variance $\sigma^2 = n/4$. To see this, observe that the degree k is the sum of n indicator variables that take on value zero or one depending whether an edge is present or not. The expected value of the sum is the sum of the expected values and the variance of the sum is the sum of the variances.

Near the mean, the binomial distribution is well approximated by the normal distribution. See Section 12.4.9 in the appendix.

$$\frac{1}{\sqrt{2\pi\sigma^2}}e^{-\frac{1}{2}\frac{(k-m)^2}{\sigma^2}} = \frac{1}{\sqrt{\pi n/2}}e^{-\frac{(k-n/2)^2}{n/2}}$$

The standard deviation of the normal distribution is $\frac{\sqrt{n}}{2}$ and essentially all of the probability mass is within an additive term $\pm c\sqrt{n}$ of the mean n/2 for some constant c and thus is certainly within a multiplicative factor of $1 \pm \varepsilon$ of n/2 for sufficiently large n.

The degree distribution of G(n, p) for general p is also binomial. Since p is the probability of an edge being present, the expected degree of a vertex is $d \approx pn$. The actual degree distribution is given by

Prob(vertex has degree k) = $\binom{n-1}{k} p^k (1-p)^{n-k-1} \approx \binom{n}{k} p^k (1-p)^{n-k}$.

The quantity $\binom{n-1}{k}$ is the number of ways of choosing k edges, out of the possible n-1 edges, and $p^k(1-p)^{n-k-1}$ is the probability that the k selected edges are present and the remaining n-k-1 are not. Since n is large, replacing n-1 by n does not cause much error.

The binomial distribution falls off exponentially fast as one moves away from the mean. However, the degree distributions of graphs that appear in many applications do not exhibit such sharp drops. Rather, the degree distributions are much broader. This is often referred to as having a "heavy tail". The term tail refers to values of a random variable far away from its mean, usually measured in number of standard deviations. Thus, although the G(n, p) model is important mathematically, more complex models are needed to represent real world graphs.

Consider an airline route graph. The graph has a wide range of degrees, from degree one or two for a small city, to degree 100, or more, for a major hub. The degree distribution is not binomial. Many large graphs that arise in various applications appear to have power law degree distributions. A power law degree distribution is one in which the number of vertices having a given degree decreases as a power of the degree, as in

Number(degree k vertices) = $c \frac{n}{k^r}$,

for some small positive real r, often just slightly less than three. Later, we will consider a random graph model giving rise to such degree distributions.

The following theorem claims that the degree distribution of the random graph G(n, p) is tightly concentrated about its expected value. That is, the probability that the degree of a vertex differs from its expected degree, np, by more than $\lambda \sqrt{np}$, drops off exponentially fast with λ .

Theorem 4.1 Let v be a vertex of the random graph G(n, p). Let α be a real number in $(0, \sqrt{np})$.

$$Prob(|np - deg(v)| \ge \alpha \sqrt{np}) \le 3e^{-\alpha^2/8}$$

Proof: The degree deg(v) is the sum of n-1 independent Bernoulli random variables, $y_1, y_2, \ldots, y_{n-1}$, where, y_i is the indicator variable that the i^{th} edge from v is present. So the theorem follows from Theorem 12.6.

Although the probability that the degree of a single vertex differs significantly from its expected value drops exponentially, the statement that the degree of every vertex is close to its expected value requires that p is $\Omega(\frac{\ln n}{n})$. That is, the expected degree grows as $\ln n$.

Corollary 4.2 Suppose ε is a positive constant. If p is $\Omega(\frac{\ln n}{n\varepsilon^2})$, then, almost surely, every vertex has degree in the range $(1 - \varepsilon)np$ to $(1 + \varepsilon)np$.

Proof: Apply Theorem 4.1 with $\alpha = \varepsilon \sqrt{np}$ to get that the probability that an individual vertex has degree outside the range $[(1 - \varepsilon)np, (1 + \varepsilon)np]$ is at most $3e^{-\varepsilon^2 np/8}$. By the union bound, the probability that some vertex has degree outside this range is at most $3ne^{-\varepsilon^2 np/8}$. For this to be o(1), it suffices for p to be $\Omega(\frac{\ln n}{n\varepsilon^2})$. Hence the Corollary.

Note that the assumption p is $\Omega(\frac{\ln n}{n\varepsilon^2})$ is necessary. If p = d/n for d a constant, for instance, then some vertices may well have degrees outside the range $[(1 - \varepsilon)d, (1 + \varepsilon)d]$. Indeed, shortly we will see that it is highly likely that for $p = \frac{1}{n}$ there is a vertex of degree $\Omega(\log n/\log \log n)$.

When p is a constant, the expected degree of vertices in G(n, p) increases with n. For example, in $G(n, \frac{1}{2})$, the expected degree of a vertex is n/2. In many real applications, we will be concerned with G(n, p) where p = d/n, for d a constant, i.e., graphs whose expected degree is a constant d independent of n. Holding d = np constant as n goes to infinity, the binomial distribution

$$\operatorname{Prob}\left(k\right) = \binom{n}{k} p^{k} \left(1-p\right)^{n-k}$$

approaches the Poisson distribution

$$Prob(k) = \frac{(np)^k}{k!}e^{-np} = \frac{d^k}{k!}e^{-d}$$

To see this, assume k = o(n) and use the approximations $n - k \cong n$, $\binom{n}{k} \cong \frac{n^k}{k!}$, and $\left(1 - \frac{d}{n}\right)^{n-k} \cong e^{-d}$ to approximate the binomial distribution by

$$\lim_{n \to \infty} \binom{n}{k} p^k (1-p)^{n-k} = \frac{n^k}{k!} \left(\frac{d}{n}\right)^k e^{-d} = \frac{d^k}{k!} e^{-d}.$$

Note that for $p = \frac{d}{n}$, where d is a constant independent of n, the probability of the binomial distribution falls off rapidly for k > d, and is essentially zero for all but some finite number of values of k. This justifies the k = o(n) assumption. Thus, the Poisson distribution is a good approximation.

Example: In $G(n, \frac{1}{n})$ many vertices are of degree one, but not all. Some are of degree zero and some are of degree greater than one. In fact, it is highly likely that there is a vertex of degree $\Omega(\log n / \log \log n)$. The probability that a given vertex is of degree k is

$$\operatorname{Prob}\left(k\right) = \binom{n}{k} \left(\frac{1}{n}\right)^{k} \left(1 - \frac{1}{n}\right)^{n-k} \approx \frac{e^{-1}}{k!}.$$

If $k = \log n / \log \log n$,

$$\log k^{k} = k \log k = \frac{\log n}{\log \log n} \left(\log \log n - \log \log \log n \right) \le \log n$$

and thus $k^k \leq n$. Since $k! \leq k^k \leq n$, the probability that a vertex has degree $k = \log n / \log \log n$ is at least $\frac{1}{k!}e^{-1} \geq \frac{1}{en}$. If the degrees of vertices were independent random variables, then this would be enough to argue that there would be a vertex of degree $\log n / \log \log n$ with probability at least $1 - (1 - \frac{1}{en})^n = 1 - e^{-\frac{1}{e}} \approx 0.31$. But the degrees are not quite independent since when an edge is added to the graph it affects the degree of two vertices. This is a minor technical point, which one can get around.

4.1.2 Existence of Triangles in G(n, d/n)

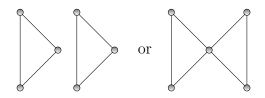
What is the expected number of triangles in $G(n, \frac{d}{n})$, when d is a constant? As the number of vertices increases one might expect the number of triangles to increase, but this is not the case. Although the number of triples of vertices grows as n^3 , the probability of an edge between two specific vertices decreases linearly with n. Thus, the probability of all three edges between the pairs of vertices in a triple of vertices being present goes down as n^{-3} , exactly canceling the rate of growth of triples.

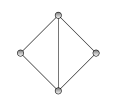
A random graph with n vertices and edge probability d/n, has an expected number of triangles that is independent of n, namely $d^3/6$. There are $\binom{n}{3}$ triples of vertices. Each triple has probability $\left(\frac{d}{n}\right)^3$ of being a triangle. Let Δ_{ijk} be the indicator variable for the triangle with vertices i, j, and k being present. That is, all three edges (i, j), (j, k), and (i, k) being present. Then the number of triangles is $x = \sum_{ijk} \Delta_{ijk}$. Even though the existence of the triangles are not statistically independent events, by linearity of expectation, which does not assume independence of the variables, the expected value of a sum of random variables is the sum of the expected values. Thus, the expected number of triangles is

$$E(x) = E\left(\sum_{ijk} \Delta_{ijk}\right) = \sum_{ijk} E(\Delta_{ijk}) = \binom{n}{3} \left(\frac{d}{n}\right)^3 \approx \frac{d^3}{6}.$$

Even though on average there are $\frac{d^3}{6}$ triangles per graph, this does not mean that with high probability a graph has a triangle. Maybe half of the graphs have $\frac{d^3}{3}$ triangles and the other half have none for an average of $\frac{d^3}{6}$ triangles. Then, with probability 1/2, a graph selected at random would have no triangle. If 1/n of the graphs had $\frac{d^3}{6}n$ triangles and the remaining graphs had no triangles, then as n goes to infinity, the probability that a graph selected at random would have a triangle would go to zero.

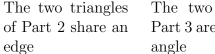
We wish to assert that with some nonzero probability there is at least one triangle in G(n, p) when $p = \frac{d}{n}$. If all the triangles were on a small number of graphs, then the number of triangles in those graphs would far exceed the expected value and hence the variance would be high. A second moment argument rules out this scenario where a small fraction of graphs have a large number of triangles and the remaining graphs have none.







The two triangles of Part 1 are either disjoint or share at most one vertex



The two triangles in Part 3 are the same triangle

Figure 4.4: The triangles in Part 1, Part 2, and Part 3 of the second moment argument for the existence of triangles in $G(n, \frac{d}{n})$.

Let's calculate $E(x^2)$ where x is the number of triangles. Write x as $x = \sum_{ijk} \Delta_{ijk}$, where Δ_{ijk} is the indicator variable of the triangle with vertices i, j, and k being present. Expanding the squared term

$$E(x^2) = E\left(\sum_{i,j,k} \Delta_{ijk}\right)^2 = E\left(\sum_{\substack{i,j,k\\i',j',k'}} \Delta_{ijk} \Delta_{i'j'k'}\right).$$

Split the above sum into three parts. In Part 1, let S_1 be the set of i, j, k and i', j', k' which share at most one vertex and hence the two triangles share no edge. In this case, Δ_{ijk} and $\Delta_{i'j'k'}$ are independent and

$$E\left(\sum_{S_1} \Delta_{ijk} \Delta_{i'j'k'}\right) = \sum_{S_1} E(\Delta_{ijk}) E(\Delta_{i'j'k'}) \le \left(\sum_{\substack{\text{all}\\ijk}} E(\Delta_{ijk})\right) \left(\sum_{\substack{\text{all}\\i'j'k'}} E(\Delta_{i'j'k'})\right) = E^2(x) E(\Delta_{ijk}) E(\Delta_{i'j'k'}) \le E(\Delta_{ijk}) E(\Delta_{i'j'k'}) = E^2(x) E(\Delta_{ijk}) E(\Delta_{ijk}) E(\Delta_{ijk}) E(\Delta_{ijk}) E(\Delta_{ijk}) = E^2(x) E(\Delta_{ijk}) E(\Delta_{ijk}$$

In Part 2, i, j, k and i', j', k' share two vertices and hence one edge. See Figure 4.4. Four vertices and five edges are involved overall. There are at most $\binom{n}{4} \in O(n^4)$, 4-vertex subsets and $\binom{4}{2}$ ways to partition the four vertices into two triangles with a common edge. The probability of all five edges in the two triangles being present is p^5 , so this part sums to $O(n^4p^5) = O(d^5/n)$ and is o(1). There are so few triangles in the graph, the probability of two triangles sharing an edge is extremely unlikely.

In Part 3, i, j, k and i', j', k' are the same sets. The contribution of this part of the summation to $E(x^2)$ is $\binom{n}{3}p^3 = \frac{d^3}{6}$. Thus, putting all three parts together, we have:

$$E(x^2) \le E^2(x) + \frac{d^3}{6} + o(1),$$

which implies

$$\operatorname{Var}(x) = E(x^2) - E^2(x) \le \frac{d^3}{6} + o(1).$$

For x to be equal to zero, it must differ from its expected value by at least its expected value. Thus,

$$\operatorname{Prob}(x=0) \le \operatorname{Prob}\left(|x-E(x)| \ge E(x)\right).$$

By Chebychev inequality,

$$\operatorname{Prob}(x=0) \le \frac{\operatorname{Var}(x)}{E^2(x)} \le \frac{d^3/6 + o(1)}{d^6/36} \le \frac{6}{d^3} + o(1).$$
(4.1)

Thus, for $d > \sqrt[3]{6} \approx 1.8$, $\operatorname{Prob}(x = 0) < 1$ and G(n, p) has a triangle with nonzero probability. For $d < \sqrt[3]{6}$ and very close to zero, there simply are not enough edges in the graph for there to be a triangle.

4.2 Phase Transitions

Many properties of random graphs undergo structural changes as the edge probability passes some threshold value. This phenomenon is similar to the abrupt phase transitions in physics, as the temperature or pressure increases. Some examples of this are the abrupt appearance of cycles in G(n, p) when p reaches 1/n and the disappearance of isolated vertices when p reaches $\frac{\log n}{n}$. The most important of these transitions is the emergence of a giant component, a connected component of size $\Theta(n)$, which happens at d = 1. Recall Figure 4.1.

Probability	Transition					
p = 0	Isolated vertices					
n = o(1)	Forest of trees, no component					
$p = o(\frac{1}{n})$	of size greater than $O(\log n)$					
$p = \frac{d}{n}, d < 1$	All components of size $O(\log n)$					
$\begin{array}{c} p = \frac{d}{n}, d < 1 \\ p = \frac{d}{n}, d = 1 \\ p = \frac{d}{n}, d > 1 \end{array}$	Components of size $O(n^{\frac{2}{3}})$					
$p = \frac{d}{n}, d > 1$	Giant component plus $O(\log n)$ components					
$p = \sqrt{2}\sqrt{\frac{\ln n}{n}}$	Diameter two					
$p = \frac{1}{2} \frac{\ln n}{n}$	Giant component plus isolated vertices					
	Disappearance of isolated vertices					
$p = \frac{\ln n}{n}$	Appearance of Hamilton circuit					
	Diameter $O(\ln n)$					
$p = \frac{1}{2}$	Clique of size $(2 - \epsilon) \log n$					

Table 1: Phase transitions

For these and many other properties of random graphs, a threshold exists where an abrupt transition from not having the property to having the property occurs. If there exists a function p(n) such that when $\lim_{n\to\infty} \frac{p_1(n)}{p(n)} = 0$, $G(n, p_1(n))$ almost surely does not

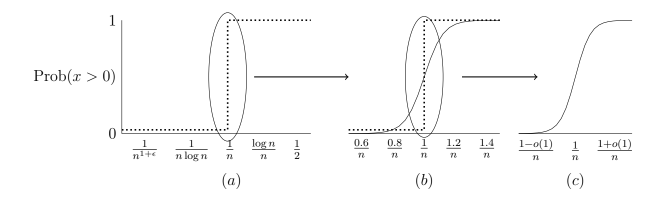


Figure 4.5: Figure 4.5(a) shows a phase transition at $p = \frac{1}{n}$. The dotted line shows an abrupt transition in $\operatorname{Prob}(x)$ from 0 to 1. For any function asymptotically less than $\frac{1}{n}$, $\operatorname{Prob}(x)>0$ is zero and for any function asymptotically greater than $\frac{1}{n}$, $\operatorname{Prob}(x)>0$ is one. Figure 4.5(b) expands the scale and shows a less abrupt change in probability unless the phase transition is sharp as illustrated by the dotted line. Figure 4.5(c) is a further expansion and the sharp transition is now more smooth.

have the property, and when $\lim_{n\to\infty} \frac{p_2(n)}{p(n)} = \infty$, $G(n, p_2(n))$ almost surely has the property, then we say that a *phase transition* occurs, and p(n) is the *threshold*. Recall that G(n, p)"almost surely does not have the property" means that the probability that it has the property goes to zero in the limit, as n goes to infinity. We shall soon see that every increasing property has a threshold. This is true not only for increasing properties of G(n, p), but for increasing properties of any combinatorial structure. If for cp(n), c < 1, the graph almost surely does not have the property and for cp(n), c > 1, the graph almost surely has the property, then p(n) is a *sharp threshold*. The existence of a giant component has a sharp threshold at 1/n. We will prove this later.

In establishing phase transitions, we often use a variable x(n) to denote the number of occurrences of an item in a random graph. If the expected value of x(n) goes to zero as n goes to infinity, then a graph picked at random almost surely has no occurrence of the item. This follows from Markov's inequality. Since x is a nonnegative random variable $\operatorname{Prob}(x \ge a) \le \frac{1}{a}E(x)$, which implies that the probability of $x(n) \ge 1$ is at most E(x(n)). That is, if the expected number of occurrences of an item in a graph goes to zero, the probability that there are one or more occurrences of the item in a randomly selected graph goes to zero. This is called the *first moment method*.

The previous section showed that the property of having a triangle has a threshold at p(n) = 1/n. If the edge probability $p_1(n)$ is o(1/n), then the expected number of triangles goes to zero and by the first moment method, the graph almost surely has no triangle. However, if the edge probability $p_2(n)$ satisfies $\frac{p_2(n)}{1/n} \to \infty$, then from (4.1), the probability of having no triangle is at most $6/d^3 + o(1) = 6/(np_2(n))^3 + o(1)$, which goes to zero. This

	At least one
	occurrence
	of item in
	10% of the
No items	graphs

$E(x) \ge 0.1$	For 10% of the
$E(x) \geq 0.1$	graphs, $x \ge 1$

Figure 4.6: If the expected fraction of the number of graphs in which an item occurs did not go to zero, then E(x), the expected number of items per graph, could not be zero. Suppose 10% of the graphs had at least one occurrence of the item. Then the expected number of occurrences per graph must be at least 0.1. Thus, $E(x) \to 0$ implies the probability that a graph has an occurrence of the item goes to zero. However, the other direction needs more work. If E(x) is large, a second moment argument is needed to conclude that the probability that a graph picked at random has an occurrence of the item is non-negligible, since there could be a large number of occurrences concentrated on a vanishingly small fraction of all graphs. The second moment argument claims that for a nonnegative random variable x with E(x) > 0, if Var(x) is $o(E^2(x))$ or alternatively if $E(x^2) \leq E^2(x) (1 + o(1))$, then almost surely x > 0.

latter case uses what we call the second moment method. The first and second moment methods are broadly used. We describe the second moment method in some generality now.

When the expected value of x(n), the number of occurrences of an item, goes to infinity, we cannot conclude that a graph picked at random will likely have a copy since the items may all appear on a vanishingly small fraction of the graphs. We resort to a technique called the *second moment method*. It is a simple idea based on Chebyshev's inequality.

Theorem 4.3 (Second Moment method) Let x(n) be a random variable with E(x) > 0. If

$$Var(x) = o\left(E^2(x)\right),$$

then x is almost surely greater than zero.

Proof: If E(x) > 0, then for x to be less than or equal to zero, it must differ from its expected value by at least its expected value. Thus,

$$\operatorname{Prob}(x \le 0) \le \operatorname{Prob}\left(|x - E(x)| \ge E(x)\right).$$

By Chebyshev inequality

$$\operatorname{Prob}\left(|x - E(x)| \ge E(x)\right) \le \frac{\operatorname{Var}(x)}{E^2(x)} \to 0.$$

Thus, $\operatorname{Prob}(x \leq 0)$ goes to zero if $\operatorname{Var}(x)$ is $o(E^2(x))$.

Corollary 4.4 Let x be a random variable with E(x) > 0. If

$$E(x^2) \le E^2(x)(1+o(1)),$$

then x is almost surely greater than zero.

Proof: If $E(x^2) \le E^2(x)(1+o(1))$, then

$$Var(x) = E(x^2) - E^2(x) \le E^2(x)o(1) = o(E^2(x)).$$

Second moment arguments are more difficult than first moment arguments since they deal with variance and without independence we do not have E(xy) = E(x)E(y). In the triangle example, dependence occurs when two triangles share a common edge. However, if $p = \frac{d}{n}$, there are so few triangles that almost surely no two triangles share a common edge and the lack of statistical independence does not affect the answer. In looking for a phase transition, almost always the transition in probability of an item being present occurs when the expected number of items transitions.

Threshold for graph diameter two (two degrees of separation)

We now present the first example of a sharp phase transition for a property. This means that slightly increasing the edge probability p near the threshold takes us from almost surely not having the property to almost surely having it. The property is that of a random graph having diameter less than or equal to two. The diameter of a graph is the maximum length of the shortest path between a pair of nodes. In other words, the property is that every pair of nodes has "at most two degrees of separation".

The following technique for deriving the threshold for a graph having diameter two is a standard method often used to determine the threshold for many other objects. Let x be a random variable for the number of objects such as triangles, isolated vertices, or Hamiltonian circuits, for which we wish to determine a threshold. Then we determine the value of p, say p_0 , where the expected value of x goes from vanishingly small to unboundedly large. For $p < p_0$ almost surely a graph selected at random will not have a copy of the item. For $p > p_0$, a second moment argument is needed to establish that the items are not concentrated on a vanishingly small fraction of the graphs and that a graph picked at random will almost surely have a copy. Our first task is to figure out what to count to determine the threshold for a graph having diameter two. A graph has diameter two if and only if for each pair of vertices iand j, either there is an edge between them or there is another vertex k to which both iand j have an edge. So, what we will count is the number of pairs i, j that fail, i.e., the number of pairs i, j that have more than two degrees of separation. The set of neighbors of i and the set of neighbors of j are random subsets of expected cardinality np. For these two sets to intersect requires $np \approx \sqrt{n}$ or $p \approx \frac{1}{\sqrt{n}}$. Such statements often go under the general name of "birthday paradox" though it is not a paradox. In what follows, we will prove a threshold of $O(\sqrt{\ln n}/\sqrt{n})$ for a graph to have diameter two. The extra factor of $\sqrt{\ln n}$ ensures that every one of the $\binom{n}{2}$ pairs of i and j has a common neighbor. When $p = c\sqrt{\frac{\ln n}{n}}$, for $c < \sqrt{2}$, the graph almost surely has diameter greater than two and for $c > \sqrt{2}$, the graph almost surely has diameter less than or equal to two.

Theorem 4.5 The property that G(n,p) has diameter two has a sharp threshold at $p = \sqrt{2}\sqrt{\frac{\ln n}{n}}$.

Proof: If G has diameter greater than two, then there exists a pair of nonadjacent vertices i and j such that no other vertex of G is adjacent to both i and j. This motivates calling such a pair *bad*.

Introduce a set of indicator random variables I_{ij} , one for each pair of vertices (i, j) with i < j, where I_{ij} is 1 if and only if the pair (i, j) is bad. Let

$$x = \sum_{i < j} I_{ij}$$

be the number of bad pairs of vertices. Putting i < j in the sum ensures each pair (i, j) is counted only once. A graph has diameter at most two if and only if it has no bad pair, i.e., x = 0. Thus, if $\lim_{n \to \infty} E(x) = 0$, then for large n, almost surely, a graph has no bad pair and hence has diameter at most two.

The probability that a given vertex is adjacent to both vertices in a pair of vertices (i, j) is p^2 . Hence, the probability that the vertex is not adjacent to both vertices is $1 - p^2$. The probability that no vertex is adjacent to the pair (i, j) is $(1 - p^2)^{n-2}$ and the probability that *i* and *j* are not adjacent is 1 - p. Since there are $\binom{n}{2}$ pairs of vertices, the expected number of bad pairs is

$$E(x) = {\binom{n}{2}} (1-p) (1-p^2)^{n-2}.$$

Setting $p = c\sqrt{\frac{\ln n}{n}}$,

$$\begin{split} E\left(x\right) &\cong \frac{n^2}{2} \left(1 - c \sqrt{\frac{\ln n}{n}}\right) \left(1 - c^2 \frac{\ln n}{n}\right)^n \\ &\cong \frac{n^2}{2} e^{-c^2 \ln n} \\ &\cong \frac{1}{2} n^{2-c^2}. \end{split}$$

For $c > \sqrt{2}$, $\lim_{n \to \infty} E(x) = 0$. Thus, by the first moment method, for $p = c\sqrt{\frac{\ln n}{n}}$ with $c > \sqrt{2}$, G(n, p) almost surely has no bad pair and hence has diameter at most two.

Next, consider the case $c < \sqrt{2}$ where $\lim_{n \to \infty} E(x) = \infty$. We appeal to a second moment argument to claim that almost surely a graph has a bad pair and thus has diameter greater than two.

$$E(x^2) = E\left(\sum_{i < j} I_{ij}\right)^2 = E\left(\sum_{i < j} I_{ij} \sum_{k < l} I_{kl}\right) = E\left(\sum_{\substack{i < j \\ k < l}} I_{ij} I_{kl}\right) = \sum_{\substack{i < j \\ k < l}} E\left(I_{ij} I_{kl}\right)$$

The summation can be partitioned into three summations depending on the number of distinct indices among i, j, k, and l. Call this number a.

$$E(x^{2}) = \sum_{\substack{i < j \\ k < l}} E(I_{ij}I_{kl}) + \sum_{\substack{i < j \\ i < k}} E(I_{ij}I_{ik}) + \sum_{\substack{i < j \\ i < k}} E(I_{ij}^{2}).$$
(4.2)

Consider the case a = 4 where i, j, k, and l are all distinct. If $I_{ij}I_{kl} = 1$, then both pairs (i, j) and (k, l) are bad and so for each $u \notin \{i, j, k, l\}$, at least one of the edges (i, u)or (j, u) is absent and, in addition, at least one of the edges (k, u) or (l, u) is absent. The probability of this for one u not in $\{i, j, k, l\}$ is $(1 - p^2)^2$. As u ranges over all the n - 4vertices not in $\{i, j, k, l\}$, these events are all independent. Thus,

$$E(I_{ij}I_{kl}) \le (1-p^2)^{2(n-4)} \le (1-c^2\frac{\ln n}{n})^{2n}(1+o(1)) \le n^{-2c^2}(1+o(1))$$

and the first sum is

$$\sum_{\substack{i < j \\ k < l}} E(I_{ij}I_{kl}) \le n^{4-2c^2}(1+o(1)).$$

For the second summation, observe that if $I_{ij}I_{ik} = 1$, then for every vertex u not equal to i, j, or k, either there is no edge between i and u or there is an edge (i, u) and both edges (j, u) and (k, u) are absent. The probability of this event for one u is

$$1 - p + p(1 - p)^2 = 1 - 2p^2 + p^3 \approx 1 - 2p^2.$$

Thus, the probability for all such u is $(1-2p^2)^{n-3}$. Substituting $c\sqrt{\frac{\ln n}{n}}$ for p yields

$$\left(1 - \frac{2c^2 \ln n}{n}\right)^{n-3} \cong e^{-2c^2 \ln n} = n^{-2c^2},$$

which is an upper bound on $E(I_{ij}I_{kl})$ for one i, j, k, and l with a = 3. Summing over all distinct triples yields n^{3-2c^2} for the second summation in (4.2).

For the third summation, since the value of I_{ij} is zero or one, $E(I_{ij}^2) = E(I_{ij})$. Thus,

$$\sum_{ij} E\left(I_{ij}^2\right) = E\left(x\right).$$

Hence, $E(x^2) \leq n^{4-2c^2} + n^{3-2c^2} + n^{2-c^2}$ and $E(x) \cong n^{2-c^2}$, from which it follows that for $c < \sqrt{2}$, $E(x^2) \leq E^2(x)(1+o(1))$. By a second moment argument, Corollary 4.4, a graph almost surely has at least one bad pair of vertices and thus has diameter greater than two. Therefore, the property that the diameter of G(n,p) is less than or equal to two has a sharp threshold at $p = \sqrt{2}\sqrt{\frac{\ln n}{n}}$

Disappearance of Isolated Vertices

The disappearance of isolated vertices in G(n,p) has a sharp threshold at $\frac{\ln n}{n}$. At this point the giant component has absorbed all the small components and with the disappearance of isolated vertices, the graph becomes connected.

Theorem 4.6 The disappearance of isolated vertices in G(n, p) has a sharp threshold of $\frac{\ln n}{n}$.

Proof: Let x be the number of isolated vertices in G(n, p). Then,

$$E(x) = n(1-p)^{n-1}.$$

Since we believe the threshold to be $\frac{\ln n}{n}$, consider $p = c \frac{\ln n}{n}$. Then,

$$\lim_{n \to \infty} E(x) = \lim_{n \to \infty} n \left(1 - \frac{c \ln n}{n} \right)^n = \lim_{n \to \infty} n e^{-c \ln n} = \lim_{n \to \infty} n^{1-c}.$$

If c > 1, the expected number of isolated vertices, goes to zero. If c < 1, the expected number of isolated vertices goes to infinity. If the expected number of isolated vertices goes to zero, it follows that almost all graphs have no isolated vertices. On the other hand, if the expected number of isolated vertices goes to infinity, a second moment argument is needed to show that almost all graphs have an isolated vertex and that the isolated vertices are not concentrated on some vanishingly small set of graphs with almost all graphs not having isolated vertices. Assume c < 1. Write $x = I_1 + I_2 + \cdots + I_n$ where I_i is the indicator variable indicating whether vertex *i* is an isolated vertex. Then $E(x^2) = \sum_{i=1}^n E(I_i^2) + 2\sum_{i < j} E(I_iI_j)$. Since I_i equals 0 or 1, $I_i^2 = I_i$ and the first sum has value E(x). Since all elements in the second sum are equal

$$E(x^{2}) = E(x) + n(n-1) E(I_{1}I_{2})$$

= $E(x) + n(n-1)(1-p)^{2(n-1)-1}$.

The minus one in the exponent 2(n-1) - 1 avoids counting the edge from vertex 1 to vertex 2 twice. Now,

$$\frac{E(x^2)}{E^2(x)} = \frac{n(1-p)^{n-1} + n(n-1)(1-p)^{2(n-1)-1}}{n^2(1-p)^{2(n-1)}}$$
$$= \frac{1}{n(1-p)^{n-1}} + (1-\frac{1}{n})\frac{1}{1-p}.$$

For $p = c \frac{\ln n}{n}$ with c < 1, $\lim_{n \to \infty} E(x) = \infty$ and

$$\lim_{n \to \infty} \frac{E(x^2)}{E^2(x)} = \lim_{n \to \infty} \left[\frac{1}{n^{1-c}} + (1 - \frac{1}{n}) \frac{1}{1 - c\frac{\ln n}{n}} \right] = 1 + o(1).$$

By the second moment argument, Corollary 4.4, the probability that x = 0 goes to zero implying that almost all graphs have an isolated vertex. Thus, $\frac{\ln n}{n}$ is a sharp threshold for the disappearance of isolated vertices. For $p = c \frac{\ln n}{n}$, when c > 1 there almost surely are no isolated vertices, and when c < 1 there almost surely are isolated vertices.

Hamilton circuits

So far in establishing phase transitions in the G(n, p) model for an item such as the disappearance of isolated vertices, we introduced a random variable x that was the number of occurrences of the item. We then determined the probability p for which the expected value of x went from zero to infinity. For values of p for which $E(x) \to 0$, we argued that with high probability, a graph generated at random had no occurrences of x. For values of x for which $E(x) \to \infty$, we used the second moment argument to conclude that with high probability, a graph generated at random had occurrences of x. That is, the occurrences that forced E(x) to infinity were not all concentrated on a vanishingly small fraction of the graphs. One might raise the question for the G(n, p) graph model, do there exist items that are so concentrated on a small fraction of the graphs that the value of p where E(x) goes from zero to infinity is not the threshold? An example where this happens is Hamilton circuits.

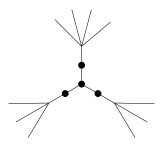


Figure 4.7: A degree three vertex with three adjacent degree two vertices. Graph cannot have a Hamilton circuit.

Let x be the number of Hamilton circuits in G(n, p) and let $p = \frac{d}{n}$ for some constant d. There are $\frac{1}{2}(n-1)!$ potential Hamilton circuits in a graph and each has probability $(\frac{d}{n})^n$ of actually being a Hamilton circuit. Thus,

$$E(x) = \frac{1}{2}(n-1)! \left(\frac{d}{n}\right)^n$$
$$\simeq \left(\frac{n}{e}\right)^n \left(\frac{d}{n}\right)^n$$
$$\to \begin{cases} 0 & d < e \\ \infty & d > e \end{cases}.$$

This suggests that the threshold for Hamilton circuits occurs when d equals Euler's constant e. This is not possible since the graph still has isolated vertices and is not even connected for $p = \frac{e}{n}$. Thus, the second moment argument is indeed necessary.

The actual threshold for Hamilton circuits is $d = \omega(\log n + \log \log n)$. For any p(n) asymptotically greater than $\frac{1}{n}(\log n + \log \log n)$, G(n, p) will have a Hamilton circuit with probability one. This is the same threshold as for the disappearance of degree one vertices. Clearly a graph with a degree one vertex cannot have a Hamilton circuit. But it may seem surprising that Hamilton circuits appear as soon as degree one vertices disappear. You may ask why at the moment degree one vertices disappear there cannot be a subgraph consisting of a degree three vertex adjacent to three degree two vertices as shown in Figure 4.7. The reason is that the frequency of degree two and three vertices in the graph is very small and the probability that four such vertices would occur together in such a subgraph is too small for it to happen with non-negligible probability.

4.3 The Giant Component

Consider G(n, p) as p grows. Starting with p = 0, the graph has n vertices and no edges. As p increases and edges are added, a forest of trees emerges. When p is o(1/n) the graph is almost surely a forest of trees, i.e., there are no cycles. When p is d/n, d a constant, cycles appear. For d < 1, no connected component has asymptotically more

than $\log n$ vertices. The number of components containing a cycle is a constant independent of n. Thus, the graph consists of a forest of trees plus a few components that have a single cycle with no $\Omega(\log n)$ size components.

At p equal to 1/n, a phase transition occurs in which a giant component emerges. The transition consists of a double jump. At p = 1/n, components of $n^{2/3}$ vertices emerge, which are almost surely trees. Then at p = d/n, d > 1, a true giant component emerges that has a number of vertices proportional to n. This is a seminal result in random graph theory and the main subject of this section. Giant components also arise in many real world graphs; the reader may want to look at large real-world graphs, like portions of the web and find the size of the largest connected component.

When one looks at the connected components of large graphs that appear in various contexts, one observes that often there is one very large component. One example is a graph formed from a data base of protein interactions¹³ where vertices correspond to proteins and edges correspond to pairs of proteins that interact. By an interaction, one means two amino acid chains that bind to each other for a function. The graph has 2735 vertices and 3602 edges. At the time we looked at the data base, the associated graph had the number of components of various sizes shown in Table ??. There are a number of small components, but only one component of size greater than 16, and that is a giant component of size 1851. As more proteins are added to the data base the giant component will grow even larger and eventually swallow up all the smaller components.

Size of component	1	2	3	4	5	6	7	8	9	10	11	12	•••	15	16	•••	1851
Number of components	48	179	50	25	14	6	4	6	1	1	1	0	0	0	1	0	1

Table 2: Size of components in the graph implicit in the database of interacting proteins.

The existence of a giant component is not unique to the graph produced from the protein data set. Take any data set that one can convert to a graph and it is likely that the graph will have a giant component, provided that the ratio of edges to vertices is a small number greater than one. Table 12.13 gives two other examples. This phenomenon, of the existence of a giant component in many real world graphs, deserves study.

Returning to G(n, p), as p increases beyond d/n, all non isolated vertices are absorbed into the giant component, and at $p = \frac{1}{2} \frac{\ln n}{n}$, the graph consists only of isolated vertices plus a giant component. At $p = \frac{\ln n}{n}$, the graph becomes completely connected. By p = 1/2, the graph is not only connected, but is sufficiently dense that it has a clique of

¹³Science 1999 July 30 Vol. 285 No. 5428 pp751-753.

ftp://ftp.cs.rochester.edu/pub/u/joel/papers.lst

Vertices are papers and edges mean that two papers shared an author.

1	2	3	4	5	6	7	8	14	27488
271	2 549	129	51	16	12	8	3	1	1

http://www.gutenberg.org/etext/3202

Vertices represent words and edges connect words that are synonyms of one another.

1	2	3	4	5	14	16	18	48	117	125	128	30242
7	1	1	1	0	1	1	1	1	1	1	1	1

Table 3: Size of components in two graphs constructed from data sets.

size $(2 - \varepsilon) \log n$ for any $\varepsilon > 0$. We prove many of these facts in this chapter.

To compute the size of a connected component of G(n, p), do a breadth first search of a component starting from an arbitrary vertex and generate an edge only when the search process needs to know if the edge exists. Start at an arbitrary vertex and mark it discovered and unexplored. At a general step, select a discovered, but unexplored vertex v, and explore it as follows. For each undiscovered vertex u, independently decide with probability p = d/n whether the edge (v, u) is in and if it is, mark u discovered and unexplored. After this, mark v explored. Discovered but unexplored vertices are called the frontier. The algorithm has found the entire connected component when the frontier becomes empty.

For each vertex u other than the start vertex, the probability that u is undiscovered after the first i steps is precisely $(1 - \frac{d}{n})^i$. A step is the full exploration of one vertex. Let z_i be the number of vertices discovered in the first i steps of the search. The distribution of z_i is Binomial $\left(n - 1, 1 - \left(1 - \frac{d}{n}\right)^i\right)$.

For large $n, 1 - (1 - \frac{d}{n})^i \cong 1 - e^{-\frac{d}{n}i}$ and the distribution of discovered vertices can be approximated by Binomial $(n, 1 - e^{-\frac{d}{n}i})$. The expected size of the set of discovered vertices is $n(1 - e^{-\frac{d}{n}i})$. The expected size of the frontier is $n(1 - e^{-\frac{d}{n}i}) - i$. Normalize the size of the frontier by dividing by n to get $1 - e^{-\frac{d}{n}i} - \frac{i}{n}$. Let $x = \frac{i}{n}$ be the normalized number of steps and let $f(x) = 1 - e^{-dx} - x$ be the normalized expected size of the frontier. When d > 1, f(0) = 0 and f'(0) = d - 1 > 0, so f is increasing at 0. But $f(1) = -e^{-d} < 0$. So, for some value $\theta, 0 < \theta < 1$, $f(\theta) = 0$. When d = 2, $\theta = 0.7968$.

For small values of i, the probability that a vertex is undiscovered after i steps is

$$\left(1-\frac{d}{n}\right)^i \approx 1-\frac{id}{n}.$$

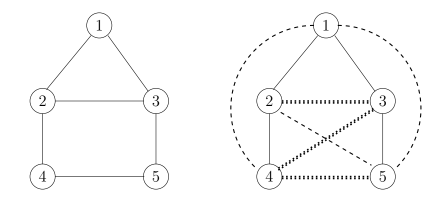


Figure 4.8: A graph (left) and the breadth first search of the graph (right). At vertex 1 the algorithm queried all edges. The solid edges are real edges, the dashed edges are edges that were queried but do not exist. At vertex 2 the algorithm queried all possible edges to vertices not yet discovered. The algorithm does not query whether the edge (2,3) exists since vertex 3 has already been discovered when the algorithm is at vertex 2. Potential edges not queried are illustrated with dotted edges.

Thus the probability that a vertex is discovered after *i* steps is approximately $\frac{id}{n}$. The expected number of discovered vertices grows as id and the expected size of the frontier grows as (d-1)i. As the fraction of discovered vertices increases, the expected rate of growth of newly discovered vertices decreases since many of the vertices adjacent to the vertex currently being searched have already been discovered. For d > 1, once $\frac{d-1}{d}n$ vertices have been discovered, the growth of newly discovered vertices slows to one at each step. Eventually for d > 1, the growth of discovering new vertices drops below one per step and the frontier starts to shrink. For d < 1, the expected size of the frontier, (d-1)i, is negative. The expected rate of growth of newly discovered vertices is less than one, even at the start.

For d > 1, the expected size of the frontier grows as (d-1)i for small *i*. The actual size of the frontier is a random variable. What is the probability that the actual size of the frontier will differ from the expected size of the frontier by a sufficient amount so that the actual size of the frontier is zero? To answer this, we need to understand the distribution of the number of discovered vertices after *i* steps. For small *i*, the probability that a vertex has been discovered is $1 - (1 - d/n)^i \approx id/n$ and the binomial distribution for the number of discovered vertices, binomial $(n, \frac{id}{n})$, is well approximated by the Poisson distribution with the same mean *id*. The probability that a total of *k* vertices have been discovered in *i* steps is approximately $e^{-di} \frac{(di)^k}{k!}$. For a connected component to have exactly *i* vertices, the frontier must drop to zero for the first time at step *i*. A necessary condition is that exactly *i* vertices must have been discovered in the first *i* steps. The probability of this is approximately

$$e^{-di}\frac{(di)^{i}}{i!} = e^{-di}\frac{d^{i}i^{i}}{i^{i}}e^{i} = e^{-(d-1)i}d^{i} = e^{-(d-1-\ln d)i}.$$

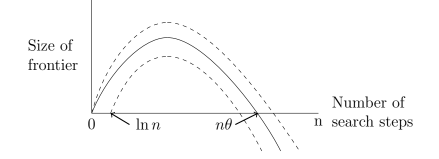


Figure 4.9: The solid curve is the expected size of the frontier. The two dashed curves indicate the high-probability range of possible values for the actual size of the frontier.

For $d \neq 1$, $d - 1 - \ln d > 0$.¹⁴ Thus the probability $e^{-(d-1-\ln d)i}$ drops off exponentially with *i*. For $i > c \ln n$ and sufficiently large *c*, the probability that the breadth first search starting from a particular vertex terminates with a component of size *i* is o(1/n) as long as the Poisson approximation is valid. In the range of this approximation, the probability that a breadth first search started from any vertex terminates with $i > c \ln n$ vertices is o(1). Intuitively, if the component has not stopped growing within $\Omega(\ln n)$ steps, it is likely to continue to grow until it becomes much larger and the expected value of the size of the frontier again becomes small. While the expected value of the frontier is large, the probability that the actual size will differ from the expected size sufficiently for the actual size of the frontier to be zero is vanishingly small.

For *i* near $n\theta$ the absolute value of the expected size of the frontier increases linearly with $|i - n\theta|$. Thus for the actual size of the frontier to be zero, z_i must deviate from its expected value by an amount proportional to $|i - n\theta|$. For values of *i* near $n\theta$, the binomial distribution can be approximated by a Gaussian distribution. The Gaussian falls off exponentially fast with the square of the distance from its mean. The distribution falls off proportional to $e^{-\frac{k^2}{\sigma^2}}$ where σ^2 is the variance and is proportional to *n*. Thus to have a non vanishing probability, *k* must be at most \sqrt{n} . This implies that the giant component is in the range $[n\theta - \sqrt{n}, n\theta + \sqrt{n}]$. Thus a component is either small or in the range $[n\theta - \sqrt{n}, n\theta + \sqrt{n}]$.

In Theorem 4.8, we prove that there is one giant component of size $\Omega(n)$ along with a number of components of size $O(\ln n)$. We first prove a technical lemma stating that the probability of a vertex being in a small component is strictly less than one and hence there is a giant component. We refer to a connected component of size $O(\log n)$ as a small component.

 $[\]overline{\int_{1^4} \text{Let } f(d) = d - 1 - \ln d. \text{ Then } \frac{\partial f}{\partial d} = 1} - \frac{1}{d} \text{ and } \frac{\partial f}{\partial d} < 0 \text{ for } d < 1 \text{ and } \frac{\partial f}{\partial d} > 0 \text{ for } d > 1. \text{ Now } f(d) = 0$ at d = 1 and is positive for all other d > 1.

Lemma 4.7 Assume d > 1. For any vertex v, the probability that cc(v), the connected component containing v, is small (i.e., of size $O(\log n)$) is a constant strictly less than 1.

Proof: Let p be the probability that cc(v) is small, i.e., the probability that the breadth first search started at v terminates before $c_1 \log n$ vertices are discovered. Slightly modify the breadth first search as follows: If in exploring a vertex u at some point, there are mundiscovered vertices, choose the number k of vertices which will be adjacent to u from the Binomial $(m, \frac{d}{n})$ distribution. Having picked k, pick one of the $\binom{m}{k}$ subsets of k undiscovered vertices to be the set of vertices adjacent to u, and make the other m-k vertices not adjacent to u. This process has the same distribution as picking each edge from uindependently at random to be present with probability d/n. As the search proceeds, m decreases. If cc(v) is small, m, the number of undiscovered vertices, is always greater than $s = n - c_1 \log n$. Modify the process once more picking k from Binomial $(s, \frac{d}{n})$ instead of from Binomial $(m, \frac{d}{n})$. Let p' be the probability that cc(v) is small for the modified process. Clearly, $p' \ge p$, so it suffices to prove that p' is a constant strictly less than one. The mean of the binomial now is $d_1 = sd/n$ which is strictly greater than one. It is clear that the probability that the modified process ends before $c_1 \log n$ vertices are discovered is at least the probability for the original process, since picking from $n - c_1 \log n$ vertices has decreased the number of newly discovered vertices each time. Modifying the process so that the newly discovered vertices are picked from a fixed size set, converts the problem to what is called a branching process. Branching processes are discussed further in Section 4.4, but we describe here everything that is needed for our analysis of the giant component.

A branching process is a method for creating a possibly infinite random tree. There is a nonnegative integer-valued random variable y that is the number of children of the node being explored. First, the root v of the tree chooses a value of y according to the distribution of y and spawns that number of children. Each of the children independently chooses a value according to the same distribution of y and spawns that many children. The process terminates when all of the vertices have spawned children. The process may go on forever. If it does terminate with a finite tree, we say that the process has become "extinct". Let Binomial $(s, \frac{d}{n})$ be the distribution of y. Let q be the probability of extinction. Then, $q \ge p'$, since, the breadth first search terminating with at most $c_1 \log n$ vertices is one way of becoming extinct. Let $p_i = {s \choose i} (d/n)^i (1 - (d/n))^{s-i}$ be the probability that y spawns i children. We have $\sum_{i=0}^{s} p_i = 1$ and $\sum_{i=1}^{s} ip_i = E(y) = ds/n > 1$.

The depth of a tree is at most the number of nodes in the tree. Let a_t be the probability that the branching process terminates at depth at most t. If the root v has no children, then the process terminates with depth one where the root is counted as a depth one node which is at most t. If v has i children, the process from v terminates at depth at most t if and only if the i sub processes, one rooted at each child of v terminate at depth t - 1 or less. The i processes are independent, so the probability that they all terminate at depth For a small number i of steps, the probability distribution of the size of the set of discovered vertices at time i is $p(k) = e^{-di} \frac{(di)^k}{k!}$ and has expected value di. Thus, the expected size of the frontier is (d-1)i. For the frontier to be empty would require that the size of the set of discovered vertices be smaller than its expected value by (d-1)i. That is, the size of the set of discovered vertices would need to be di - (d-1)i = i. The probability of this is

$$e^{-di}\frac{(di)^{i}}{i!} = e^{-di}\frac{d^{i}i^{i}}{i^{i}}e^{i} = e^{-(d-1)i}d^{i} = e^{-(d-1-\ln d)i}$$

which drops off exponentially fast with *i* provided d > 1. Since $d - 1 - \ln d$ is some constant c > 0, the probability is e^{-ci} which for $i = \ln n$ is $e^{-c \ln n} = \frac{1}{n^c}$. Thus, with high probability, the largest small component in the graph is of size at most $\ln n$.

Illustration 4.1

at most t-1 is exactly a_{t-1}^i . With this we get:

$$a_t = p_0 + \sum_{i=1}^{s} p_i a_{t-1}^i = \sum_{i=0}^{s} p_i a_{t-1}^i.$$

We have $a_1 = p_0 < 1$. There is a constant $\alpha \in [p_0, 1)$ such that whenever $a_{t-1} \leq \alpha$, the above recursion implies that $a_t \leq \alpha$. This would finish the proof since then $a_1 \leq \alpha$ implies $a_2 \leq \alpha$ which implies $a_3 \leq \alpha$ etc. and so $q = \lim_{t \to \infty} a_t \leq \alpha$.

To prove the claim, consider the polynomial

$$h(x) = x - \sum_{i=0}^{s} p_i x^i.$$

We see that h(1) = 0 and $h'(1) = 1 - \sum_{i=1}^{s} ip_i \approx 1 - \frac{sd}{n}$, which is at most a strictly negative constant. By continuity of $h(\cdot)$, there exists some $x_0 < 1$ such that $h(x) \ge 0$ for $x \in [x_0, 1]$. Take $\alpha = \operatorname{Max}(x_0, p_0)$. Now since $\sum_{i=0}^{s} p_i x^i$ has all nonnegative coefficients, it is an increasing function of x and so if a_{t-1} is at least α , then, $\sum_{i=0}^{s} p_i a_{t-1}^i$ is at least $\sum_{i=0}^{s} p_i \alpha^i \ge \alpha$. Now, if $a_{t-1} \le \alpha$,

$$a_t = \sum_{i=0}^s p_i a_{t-1}^i \ge \sum_{i=1}^s p_i \alpha^i = \alpha - h(\alpha) \le \alpha,$$

proving the claim.

We now prove in Theorem 4.8 that in $G(n, \frac{d}{n})$, d > 1, there is one giant component containing a fraction of the *n* vertices and that the remaining vertices are in components of size less than some constant c_1 times $\log n$. There are no components greater than $c_1 \log n$ other than the giant component. **Theorem 4.8** Let p=d/n with d > 1.

- 1. There are constants c_1 and c_2 such that the probability that there is a connected component of size between $c_1 \log n$ and $c_2 n$ is at most 1/n.
- 2. The number of vertices in components of size $O(\ln n)$ is almost surely at most cn for some c < 1. Thus, with probability 1 - o(1), there is a connected component of size $\Omega(n)$.
- 3. The probability that there are two or more connected components, each of size more than $n^{2/3}$, is at most 1/n.

Proof: We begin with (1). If a particular vertex v is in a connected component of size k, then the frontier of the breadth first search process started at v becomes empty for the first time at time k implying that $z_k = k$. Now z_k has distribution $\text{Binomial}(n-1, 1-(1-(d/n))^k)$. Let $c_2 = \frac{d-1}{d^2}$ and assume $k < c_2 n$ and $k > c_1 \log n$ for a sufficiently large c_1 . By truncated Taylor series expansion, it is easy to see that

$$(1 - \frac{d}{n})^k \le 1 - \frac{kd}{n} + \frac{k^2d^2}{2n^2}.$$

Thus, $E(z_k) \ge kd - (k^2d^2/2n)$. So, if z_k were to be exactly k, then its deviation from its mean is at least $kd - (k^2d^2/(2n)) - k \ge (k(d-1)/2) \ge \Omega(k)$. So by a Chernoff bound $\operatorname{Prob}(z_k = k) \le e^{-\Omega(k)}$. If k is greater than $c_1 \log n$ for some sufficiently large c_1 , then $z_k = k$ with probability at most $1/n^3$ and this along with a union bound gives (1).

Next consider (2). For a vertex v, let cc(v) denote the set of vertices in the connected component containing v. By (1), almost surely, cc(v) is a small set of size at most $c_1 \log n$ or a large set of size at least c_2n for every vertex v. The central part of the proof of (2) that the probability of a vertex being in a small component is strictly less than one was established in Lemma 4.7. Let x be the number of vertices in a small connected component. Lemma 4.7 implies that the expectation of the random variable x equals the number of vertices in small connected components is at most some c_3n , for a constant c_3 strictly less than one. But we need to show that for any graph almost surely the actual number x of such vertices is at most some constant strictly less than one times n. For this, we use the second moment method. In this case, the proof that the variance of x is $o(E^2(x))$ is easy. Let x_i be the indicator random variable of the event that cc(i) is small. Let S and T run over all small sets. Noting that for $i \neq j$, cc(i) and cc(j) either are the same or are disjoint,

$$\begin{split} E(x^2) &= E\left(\left(\sum_{i=1}^n x_i\right)^2\right) = \sum_{i,j} E(x_i x_j) = \sum_i E(x_i^2) + \sum_{i \neq j} E(x_i x_j) \\ &= E(x) + \sum_{i \neq j} \sum_S \operatorname{Prob}\left(\operatorname{cc}(i) = \operatorname{cc}(j) = S\right) + \sum_{i \neq j} \sum_{\substack{S.T. \\ \text{disjoint}}} \operatorname{Prob}\left(\operatorname{cc}(i) = S; \operatorname{cc}(j) = T\right) \\ &= E(x) + \sum_{i \neq j} \sum_{\substack{S:i,j \in S}} \operatorname{Prob}(S \text{ is a c.c.}) + \sum_{i \neq j} \sum_{\substack{S.T. : i \in S. j \in T \\ \text{disjoint}}} \operatorname{Prob}(S, T \text{ each a cc}) \\ &= E(x) + \sum_{i \neq j} \sum_{\substack{S:i,j \in S}} \operatorname{Prob}\left(S \text{ is a c.c.}\right) \\ &+ \sum_{i \neq j} \sum_{\substack{S:T. : i \in S. j \in T \\ \text{disjoint}}} \operatorname{Prob}\left(S \text{ is a c.c.}\right) \operatorname{Prob}\left(T \text{ is a c.c.}\right)(1 - p)^{-|S||T} \\ &\leq O(n) + (1 - p)^{-|S||T|} \left(\sum_S \operatorname{Prob}\left(\operatorname{cc}(i) = S\right)\right) \left(\sum_T \operatorname{Prob}\left(\operatorname{cc}(j) = T\right)\right) \\ &\leq O(n) + (1 + o(1)) E(x)E(x). \end{split}$$

In the next to last line, if S containing i and T containing j are disjoint sets, then the two events, S is a connected component and T is a connected component, depend on disjoint sets of edges except for the |S||T| edges between S vertices and T vertices. Let c_4 be a constant in the interval $(c_3, 1)$. Then, by Chebyshev inequality,

$$\operatorname{Prob}(x > c_4 n) \le \frac{\operatorname{Var}(x)}{(c_4 - c_3)^2 n^2} \le \frac{O(n) + o(1)c_3^2 n^2}{(c_4 - c_3)^2 n^2} = o(1).$$

For the proof of (3) suppose a pair of vertices u and v belong to two different connected components, each of size at least $n^{2/3}$. We show that with high probability, they should have merged into one component producing a contradiction. First, run the breadth first search process starting at v for $\frac{1}{2}n^{2/3}$ steps. Since v is in a connected component of size $n^{2/3}$, there are $\Omega(n^{2/3})$ frontier vertices. The expected size of the frontier continues to grow until some constant times n and the actual size of the frontier does not differ significantly from the expected size. The size of the component also grows linearly with n. Thus, the frontier is of size $n^{\frac{2}{3}}$. See Exercise 4.22. By the assumption, u does not belong to this connected component. Now, temporarily stop the breadth first search tree of v and begin a breadth first search tree starting at u, again for $\frac{1}{2}n^{2/3}$ steps. It is important to understand that this change of order of building G(n, p) does not change the resulting graph. We can choose edges in any order since the order does not affect independence or conditioning. The breadth first search tree from u also will have $\Omega(n^{2/3})$ frontier vertices with high probability. Now grow the u tree further. The probability that none of the edges between the two frontier sets is encountered is $(1-p)^{\Omega(n^{4/3})} \leq e^{-\Omega(dn^{1/3})}$, which converges to zero. So almost surely, one of the edges is encountered and u and v end up in the same connected component. This argument shows for a particular pair of vertices u and v, the probability that they belong to different large connected components is very small. Now use the union bound to conclude that this does not happen for any of the $\binom{n}{2}$ pairs of vertices. The details are left to the reader.

For the d < 1 case, almost surely, there is no connected component of size $\Omega(\ln n)$.

Theorem 4.9 Let p=d/n with d < 1. The probability that G(n,p) has a component of size more than $c\frac{\ln n}{(1-d)^2}$ is at most 1/n for a suitable constant c depending on d but not on n.

Proof: There is a connected component of size at least k containing a particular vertex v only if the breadth first search started at v has a nonempty frontier at all times up to k. Let z_k be the number of discovered vertices after k steps. The probability that v is in a connected component of size greater than or equal to k is less than or equal to $\operatorname{Prob}(z_k > k)$. Now the distribution of z_k is $\operatorname{Binomial}(n-1, 1-(1-d/n)^k)$. Since $(1-d/n)^k \ge 1 - dk/n$, the mean of $\operatorname{Binomial}(n-1, 1-(1-d/n)^k)$ is less than the mean of $\operatorname{Binomial}(n, \frac{dk}{n})$. Since $\operatorname{Binomial}(n, \frac{dk}{n})$ has mean dk, the mean of z_k is at most dk where d < 1. By a Chernoff bound, the probability that z_k is greater than k is at most e^{-c_0k} for some constant $c_0 > 0$. If $k \ge c \ln n$ for a suitably large c, then this probability is at most $1/n^2$. This bound is for a single vertex v. Multiplying by n for the union bound completes the proof.

4.4 Branching Processes

As discussed in the previous section, a *branching process* is a method for creating a random tree. Starting with the root node, each node has a probability distribution for the number of its children. The root of the tree denotes a parent and its descendants are the children with their descendants being the grandchildren. The children of the root are the first generation, their children the second generation, and so on. Branching processes have obvious applications in population studies, and as we saw earlier in exploring a connected component in a random graph. Here, we give a more in-depth discussion of their properties.

We analyze a simple case of a branching process where the distribution of the number of children at each node in the tree is the same. The basic question asked is what is the probability that the tree is finite, i.e., the probability that the branching process dies out? This is called the *extinction probability*.

Our analysis of the branching process will give the probability of extinction, as well as the expected size of the components conditioned on extinction. This will imply that in $G(n, \frac{d}{n})$, with d > 1, there is one giant component of size $\Omega(n)$, the rest of the components are $O(\ln n)$ in size and the expected size of the small components is O(1).

An important tool in our analysis of branching processes is the generating function. The generating function for a nonnegative integer valued random variable y is $f(x) = \sum_{i=0}^{\infty} p_i x^i$ where p_i is the probability that y equals i. The reader not familiar with generating functions should consult Section 12.8 of the appendix.

Let the random variable z_j be the number of children in the j^{th} generation and let $f_j(x)$ be the generating function for z_j . Then $f_1(x) = f(x)$ is the generating function for the first generation where f(x) is the generating function for the number of children at a node in the tree. The generating function for the 2^{nd} generation is $f_2(x) = f(f(x))$. In general, the generating function for the $j + 1^{st}$ generation is given by $f_{j+1}(x) = f_j(f(x))$. To see this, observe two things.

First, the generating function for the sum of two identically distributed integer valued random variables x_1 and x_2 is the square of their generating function

$$f^{2}(x) = p_{0}^{2} + (p_{0}p_{1} + p_{1}p_{0})x + (p_{0}p_{2} + p_{1}p_{1} + p_{2}p_{0})x^{2} + \cdots$$

For $x_1 + x_2$ to have value zero, both x_1 and x_2 must have value zero, for $x_1 + x_2$ to have value one, exactly one of x_1 or x_2 must have value zero and the other have value one, and so on. In general, the generating function for the sum of *i* independent random variables, each with generating function f(x), is $f^i(x)$.

The second observation is that the coefficient of x^i in $f_j(x)$ is the probability of there being *i* children in the j^{th} generation. If there are *i* children in the j^{th} generation, the number of children in the $j + 1^{st}$ generation is the sum of *i* independent random variables each with generating function f(x). Thus, the generating function for the $j + 1^{st}$ generation, given *i* children in the j^{th} generation, is $f^i(x)$. The generating function for the $j + 1^{st}$ generation is given by

$$f_{j+1}(x) = \sum_{i=0}^{\infty} \operatorname{Prob}(z_j = i) f^i(x).$$

If $f_j(x) = \sum_{i=0}^{\infty} a_i x^i$, then f_{j+1} is obtained by substituting f(x) for x in $f_j(x)$.

Since f(x) and its iterates, f_2, f_3, \ldots , are all polynomials in x with nonnegative coefficients, f(x) and its iterates are all monotonically increasing and convex on the unit interval. Since the probabilities of the number of children of a node sum to one, if $p_0 < 1$, some coefficient of x to a power other than zero in f(x) is nonzero and f(x) is strictly increasing.

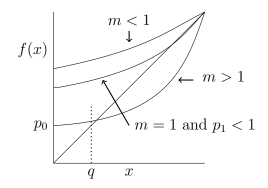


Figure 4.10: Illustration of the root of equation f(x) = x in the interval [0,1).

Let q be the probability that the branching process dies out. If there are *i* children in the first generation, then each of the *i* subtrees must die out and this occurs with probability q^i . Thus, q equals the summation over all values of *i* of the product of the probability of *i* children times the probability that *i* subtrees will die out. This gives $q = \sum_{i=0}^{\infty} p_i q^i$. Thus, q is the root of $x = \sum_{i=0}^{\infty} p_i x^i$, that is x = f(x).

This suggests focusing on roots of the equation f(x) = x in the interval [0,1]. The value x = 1 is always a root of the equation f(x) = x since $f(1) = \sum_{i=0}^{\infty} p_i = 1$. When is there a smaller nonnegative root? The derivative of f(x) at x = 1 is $f'(1) = p_1 + 2p_2 + 3p_3 + \cdots$. Let m = f'(1). Thus, m is the expected number of children of a node. If m > 1, one might expect the tree to grow forever, since each node at time j is expected to have more than one child. But this does not imply that the probability of extinction is zero. In fact, if $p_0 > 0$, then with positive probability, the root will have no children and the process will become extinct right away. Recall that for $G(n, \frac{d}{n})$, the expected number of children is d, so the parameter m plays the role of d.

If m < 1, then the slope of f(x) at x = 1 is less than one. This fact along with convexity of f(x) implies that f(x) > x for x in [0, 1) and there is no root of f(x) = x in the interval [0, 1).

If m = 1 and $p_1 < 1$, then once again convexity implies that f(x) > x for $x \in [0, 1)$ and there is no root of f(x) = x in the interval [0, 1). If m = 1 and $p_1 = 1$, then f(x) is the straight line f(x) = x.

If m > 1, then the slope of f(x) is greater than the slope of x at x = 1. This fact, along with convexity of f(x), implies f(x) = x has a unique root in [0,1). When $p_0 = 0$, the root is at x = 0.

Let q be the smallest nonnegative root of the equation f(x) = x. For m < 1 and for

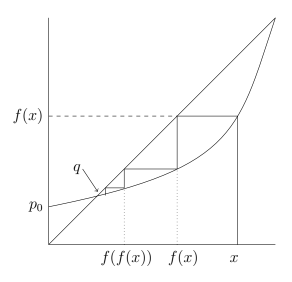


Figure 4.11: Illustration of convergence of the sequence of iterations $f_1(x), f_2(x), \ldots$ to q.

m=1 and $p_0 < 1$, q equals one and for m > 1, q is strictly less than one. We shall see that the value of q is the *extinction probability* of the branching process and that 1-q is the *immortality probability*. That is, q is the probability that for some j, the number of children in the j^{th} generation is zero. To see this, note that for m > 1, $\lim_{j \to \infty} f_j(x) = q$ for $0 \le x < 1$. Figure 4.11 illustrates the proof which is given in Lemma 4.10. Similarly note that when m < 1 or m = 1 with $p_0 < 1$, $f_j(x)$ approaches one as j approaches infinity.

Lemma 4.10 Assume m > 1. Let q be the unique root of f(x)=x in [0,1). In the limit as j goes to infinity, $f_j(x) = q$ for x in [0,1).

Proof: If $0 \le x \le q$, then $x < f(x) \le f(q)$ and iterating this inequality

$$x < f_1(x) < f_2(x) < \dots < f_j(x) < f(q) = q$$

Clearly, the sequence converges and it must converge to a fixed point where f(x) = x. Similarly, if $q \le x < 1$, then $f(q) \le f(x) < x$ and iterating this inequality

$$x > f_1(x) > f_2(x) > \dots > f_j(x) > f(q) = q.$$

In the limit as j goes to infinity $f_j(x) = q$ for all $x, 0 \le x < 1$.

Recall that $f_j(x)$ is the generating function $\sum_{i=0}^{\infty} \operatorname{Prob}(z_j = i) x^i$. The fact that in the limit the generating function equals the constant q, and is not a function of x, says that $\operatorname{Prob}(z_j = 0) = q$ and $\operatorname{Prob}(z_j = i) = 0$ for all finite nonzero values of i. The remaining probability is the probability of a nonfinite component. Thus, when m > 1, q is the extinction probability and 1-q is the probability that z_j grows without bound, i.e., immortality.

Theorem 4.11 Consider a tree generated by a branching process. Let f(x) be the generating function for the number of children at each node.

- 1. If the expected number of children at each node is less than or equal to one, then the probability of extinction is one unless the probability of exactly one child is one.
- 2. If the expected number of children of each node is greater than one, then the probability of extinction is the unique solution to f(x) = x in [0, 1).

Proof: Let p_i be the probability of *i* children at each node. Then $f(x) = p_0 + p_1 x + p_2 x^2 + \cdots$ is the generating function for the number of children at each node and $f'(1) = p_1 + 2p_2 + 3p_3 + \cdots$ is the slope of f(x) at x = 1. Observe that f'(1) is the expected number of children at each node.

Since the expected number of children at each node is the slope of f(x) at x = 1, if the expected number of children is less than or equal to one, the slope of f(x) at x = 1is less than or equal to one and the unique root of f(x) = x in (0,1] is at x = 1 and the probability of extinction is one unless f'(1) = 1 and $p_1 = 1$. If f'(1) = 1 and $p_1 = 1$, f(x) = x and the tree is an infinite degree one chain. If the slope of f(x) at x = 1 is greater than one, then the probability of extinction is the unique solution to f(x) = x in [0, 1).

A branching process with m < 1 or m=1 and $p_1 < 1$ dies out with probability one. If m=1 and $p_1 = 1$, then the branching process consists of an infinite chain with no fan out. If m > 1, then the branching process will die out with some probability less than one unless $p_0 = 0$ in which case it cannot die out, since a node always has at least one descendent.

Note that the branching process corresponds to finding the size of a component in an infinite graph. In a finite graph, the probability distribution of descendants is not a constant as more and more vertices of the graph get discovered.

The simple branching process defined here either dies out or goes to infinity. In biological systems there are other factors, since processes often go to stable populations. One possibility is that the probability distribution for the number of descendants of a child depends on the total population of the current generation.

Expected size of extinct families

We now show that the expected size of an extinct family is finite, provided that $m \neq 1$. Note that at extinction, the size must be finite. However, the expected size at extinction could conceivably be infinite, if the probability of dying out did not decay fast enough. To see how the expected value of a random variable that is always finite could be infinite, let x be an integer valued random variable. Let p_i be the probability that x = i. If $\sum_{i=1}^{\infty} p_i = 1$, then with probability one, x will be finite. However, the expected value of x may be infinite. That is, $\sum_{i=0}^{\infty} ip_i = \infty$. For example, if for i > 0, $p_i = \frac{6}{\pi^2} \frac{1}{i^2}$, then $\sum_{i=1}^{\infty} p_i = 1$, but $\sum_{i=1}^{\infty} ip_i = \infty$. The value of the random variable x is always finite, but its expected value is infinite. This does not happen in a branching process, except in the special case where the slope m = f'(1) equals one and $p_1 \neq 1$

Lemma 4.12 If the slope m = f'(1) does not equal one, then the expected size of an extinct family is finite. If the slope m equals one and $p_1 = 1$, then the tree is an infinite degree one chain and there are no extinct families. If m=1 and $p_1 < 1$, then the expected size of the extinct family is infinite.

Proof: Let z_i be the random variable denoting the size of the i^{th} generation and let q be the probability of extinction. The probability of extinction for a tree with k children in the first generation is q^k since each of the k children has an extinction probability of q. Note that the expected size of z_1 , the first generation, over extinct trees will be smaller than the expected size of z_1 over all trees since when the root node has a larger number of children than average, the tree is more likely to be infinite.

By Bayes rule

$$\operatorname{Prob}\left(z_{1}=k|\operatorname{extinction}\right)=\operatorname{Prob}\left(z_{1}=k\right)\frac{\operatorname{Prob}\left(\operatorname{extinction}|z_{1}=k\right)}{\operatorname{Prob}\left(\operatorname{extinction}\right)}=p_{k}\frac{q^{k}}{q}=p_{k}q^{k-1}.$$

Knowing the probability distribution of z_1 given extinction, allows us to calculate the expected size of z_1 given extinction.

$$E(z_1|\text{extinction}) = \sum_{k=0}^{\infty} k p_k q^{k-1} = f'(q).$$

We now prove, using independence, that the expected size of the i^{th} generation given extinction is

$$E(z_i|\text{extinction}) = \left(f'(q)\right)^i.$$

For $i = 2, z_2$ is the sum of z_1 independent random variables, each independent of the random variable z_1 . So, $E(z_2|z_1 = j$ and extinction) = E(sum of j copies of z_1 extinction) = $jE(z_1|$ extinction). Summing over all values of j

$$E(z_2|\text{extinction}) = \sum_{j=1}^{\infty} E(z_2|z_1 = j \text{ and extinction}) \text{Prob}(z_1 = j|\text{extinction})$$
$$= \sum_{j=1}^{\infty} j E(z_1|\text{extinction}) \text{Prob}(z_1 = j|\text{extinction})$$
$$= E(z_1|\text{extinction}) \sum_{j=1}^{\infty} j \text{Prob}(z_1 = j|\text{extinction}) = E^2(z_1|\text{extinction}).$$

Since $E(z_1|\text{extinction}) = f'(q)$, $E(z_2|\text{extinction}) = (f'(q))^2$. Similarly, $E(z_i|\text{extinction}) = (f'(q))^i$. The expected size of the tree is the sum of the expected sizes of each generation. That is,

Expected size of
tree given extinction
$$=\sum_{i=0}^{\infty} E(z_i | \text{extinction}) = \sum_{i=0}^{\infty} (f'(q))^i = \frac{1}{1 - f'(q)}$$

Thus, the expected size of an extinct family is finite since f'(q) < 1 provided $m \neq 1$.

The fact that f'(q) < 1 is illustrated in Figure 4.10. If m < 1, then q=1 and f'(q) = m is less than one. If m > 1, then $q \in [0, 1)$ and again f'(q) < 1 since q is the solution to f(x) = x and f'(q) must be less than one for the curve f(x) to cross the line x. Thus, for m < 1 or m > 1, f'(q) < 1 and the expected tree size of $\frac{1}{1-f'(q)}$ is finite. For m=1 and $p_1 < 1$, one has q=1 and thus f'(q) = 1 and the formula for the expected size of the tree diverges.

4.5 Cycles and Full Connectivity

This section considers when cycles form and when the graph becomes fully connected. For both of these problems, we look at each subset of k vertices and see when they form either a cycle or when they form a connected component.

4.5.1 Emergence of Cycles

The emergence of cycles in G(n, p) has a threshold when p equals to 1/n. However, the threshold is not sharp.

Theorem 4.13 The threshold for the existence of cycles in G(n,p) is p = 1/n.

Proof: Let x be the number of cycles in G(n, p). To form a cycle of length k, the vertices can be selected in $\binom{n}{k}$ ways. Given the k vertices of the cycle, they can be ordered by arbitrarily selecting a first vertex, then a second vertex in one of k-1 ways, a third in one of k-2 ways, etc. Since a cycle and its reversal are the same cycle, divide by 2. Thus, there are $\binom{n}{k} \frac{(k-1)!}{2}$ possible cycles of length k and

$$E(x) = \sum_{k=3}^{n} \binom{n}{k} \frac{(k-1)!}{2} p^{k} \le \sum_{k=3}^{n} \frac{n^{k}}{2k} p^{k} \le \sum_{k=3}^{n} (np)^{k} = (np)^{3} \frac{1-(np)^{n-2}}{1-np} \le 2(np)^{3},$$

provided that np < 1/2. When p is asymptotically less than 1/n, then $\lim_{n \to \infty} np = 0$ and $\lim_{n \to \infty} \sum_{k=3}^{n} (np)^k = 0$. So, as n goes to infinity, E(x) goes to zero. Thus, the graph almost surely has no cycles by the first moment method. A second moment argument can be used to show that for p = d/n, d > 1, a graph will have a cycle with probability tending to one.

Property	Threshold
cycles	1/n
giant component	1/n
giant component + isolated vertices	$\frac{1}{2}\frac{\ln n}{n}$
connectivity, disappearance of isolated vertices	$\frac{\ln n}{n}$
diameter two	$\sqrt{\frac{2\ln n}{n}}$

The argument above does not yield a sharp threshold since we argued that $E(x) \to 0$ only under the assumption that p is asymptotically less than $\frac{1}{n}$. A sharp threshold requires $E(x) \to 0$ for p = d/n, d < 1.

Consider what happens in more detail when p = d/n, d a constant.

$$E(x) = \sum_{k=3}^{n} {\binom{n}{k}} \frac{(k-1)!}{2} p^{k}$$

= $\frac{1}{2} \sum_{k=3}^{n} \frac{n(n-1)\cdots(n-k+1)}{k!} (k-1)! p^{k}$
= $\frac{1}{2} \sum_{k=3}^{n} \frac{n(n-1)\cdots(n-k+1)}{n^{k}} \frac{d^{k}}{k}.$

E(x) converges if d < 1, and diverges if $d \ge 1$. If d < 1, $E(x) \le \frac{1}{2} \sum_{k=3}^{n} \frac{d^{k}}{k}$ and $\lim_{n \to \infty} E(x)$ equals a constant greater than zero. If d = 1, $E(x) = \frac{1}{2} \sum_{k=3}^{n} \frac{n(n-1)\cdots(n-k+1)}{n^{k}} \frac{1}{k}$. Consider only the first log *n* terms of the sum. Since $\frac{n}{n-i} = 1 + \frac{i}{n-i} \le e^{i/n-i}$, it follows that $\frac{n(n-1)\cdots(n-k+1)}{n^{k}} \ge 1/2$. Thus,

$$E(x) \ge \frac{1}{2} \sum_{k=3}^{\log n} \frac{n(n-1)\cdots(n-k+1)}{n^k} \frac{1}{k} \ge \frac{1}{4} \sum_{k=3}^{\log n} \frac{1}{k}.$$

Then, in the limit as n goes to infinity

$$\lim_{n \to \infty} E(x) \ge \lim_{n \to \infty} \frac{1}{4} \sum_{k=3}^{\log n} \frac{1}{k} \ge \lim_{n \to \infty} (\log \log n) = \infty.$$

For p = d/n, d < 1, E(x) converges to a nonzero constant. For d > 1, E(x) converges to infinity and a second moment argument shows that graphs will have an unbounded number of cycles increasing with n.

4.5.2 Full Connectivity

As p increases from p = 0, small components form. At p = 1/n a giant component emerges and swallows up smaller components, starting with the larger components and ending up swallowing isolated vertices forming a single connected component at $p = \frac{\ln n}{n}$, at which point the graph becomes connected. We begin our development with a technical lemma.

Lemma 4.14 The expected number of connected components of size k in G(n, p) is at most

$$\binom{n}{k}k^{k-2}p^{k-1}(1-p)^{kn-k^2}.$$

Proof: The probability that k vertices form a connected component consists of the product of two probabilities. The first is the probability that the k vertices are connected, and the second is the probability that there are no edges out of the component to the remainder of the graph. The first probability is at most the sum over all spanning trees of the k vertices, that the edges of the spanning tree are present. The "at most" in the lemma statement is because G(n, p) may contain more than one spanning tree on these nodes and, in this case, the union bound is higher than the actual probability. There are k^{k-2} spanning trees of k nodes. See Section 12.9.6 in the appendix. The probability that there are no edges connecting the k vertices to the remainder of the graph is $(1-p)^{k(n-k)}$. Thus, the probability of one particular set of k vertices forming a connected component is at most $\binom{n}{k}k^{k-2}p^{k-1}(1-p)^{kn-k^2}$.

We now prove that for $p = \frac{1}{2} \frac{\ln n}{n}$, the giant component has absorbed all small components except for isolated vertices.

Theorem 4.15 For $p = c \frac{\ln n}{n}$ with c > 1/2, almost surely there are only isolated vertices and a giant component. For c > 1, almost surely the graph is connected.

Proof: We prove that almost surely for c > 1/2, there is no connected component with k vertices for any $k, 2 \le k \le n/2$. This proves the first statement of the theorem since, if there were two or more components that are not isolated vertices, both of them could not be of size greater than n/2. The second statement that for c > 1 the graph is connected then follows from Theorem 4.6 which states that isolated vertices disappear at c = 1.

We now show that for $p = c \frac{\ln n}{n}$, the expected number of components of size k, $2 \le k \le n/2$, is less than n^{1-2c} and thus for c > 1/2 there are no components, except for isolated vertices and the giant component. Let x_k be the number of connected components of size k. Substitute $p = c \frac{\ln n}{n}$ into $\binom{n}{k} k^{k-2} p^{k-1} (1-p)^{kn-k^2}$ and simplify using $\binom{n}{k} \le (en/k)^k$, $1-p \le e^{-p}$, k-1 < k, and $x = e^{\ln x}$ to get

$$E(x_k) \le \exp\left(\ln n + k + k\ln\ln n - 2\ln k + k\ln c - ck\ln n + ck^2\frac{\ln n}{n}\right)$$

Keep in mind that the leading terms here for large k are the last two and, in fact, at k = n, they cancel each other so that our argument does not prove the fallacious statement for $c \ge 1$ that there is no connected component of size n, since there is. Let

$$f(k) = \ln n + k + k \ln \ln n - 2 \ln k + k \ln c - ck \ln n + ck^2 \frac{\ln n}{n}.$$

Differentiating with respect to k,

$$f'(k) = 1 + \ln \ln n - \frac{2}{k} + \ln c - c \ln n + \frac{2ck \ln n}{n}$$

and

$$f''(k) = \frac{2}{k^2} + \frac{2c\ln n}{n} > 0.$$

Thus, the function f(k) attains its maximum over the range [2, n/2] at one of the extreme points 2 or n/2. At k = 2, $f(2) \approx (1 - 2c) \ln n$ and at k = n/2, $f(n/2) \approx -c\frac{n}{4} \ln n$. So f(k) is maximum at k = 2. For k = 2, $E(x_k) = e^{f(k)}$ is approximately $e^{(1-2c)\ln n} = n^{1-2c}$ and is geometrically falling as k increases from 2. At some point $E(x_k)$ starts to increase but never gets above $n^{-\frac{c}{4}n}$. Thus, the expected sum of the number of components of size k, for $2 \le k \le n/2$ is

$$E\left(\sum_{k=2}^{n/2} x_k\right) = O(n^{1-2c}).$$

This expected number goes to zero for c > 1/2 and the first-moment method implies that, almost surely, there are no components of size between 2 and n/2. This completes the proof of Theorem 4.15.

4.5.3 Threshold for $O(\ln n)$ Diameter

We now show that within a constant factor of the threshold for graph connectivity, not only is the graph connected, but its diameter is $O(\ln n)$. That is, if p is $\Omega(\ln n/n)$, the diameter of G(n, p) is $O(\ln n)$.

Consider a particular vertex v. Let S_i be the set of vertices at distance i from v. We argue that as i grows, $|S_1| + |S_2| + \cdots + |S_i|$ grows by a constant factor up to a size of n/1000. This implies that in $O(\ln n)$ steps, at least n/1000 vertices are connected to v. Then, there is a simple argument at the end of the proof of Theorem 4.17 that a pair of n/1000 sized subsets, connected to two different vertices v and w, have an edge between them.

Lemma 4.16 Consider G(n,p) for sufficiently large n with $p = c \ln n/n$ for any c > 0. Let S_i be the set of vertices at distance i from some fixed vertex v. If $|S_1| + |S_2| + \cdots + |S_i| \le n/1000$, then

$$Prob\left(|S_{i+1}| < 2(|S_1| + |S_2| + \dots + |S_i|)\right) \le e^{-10|S_i|}.$$

Proof: Let $|S_i| = k$. For each vertex u not in $S_1 \cup S_2 \cup \ldots \cup S_i$, the probability that u is not in S_{i+1} is $(1-p)^k$ and these events are independent. So, $|S_{i+1}|$ is the sum of $n - (|S_1| + |S_2| + \cdots + |S_i|)$ independent Bernoulli random variables, each with probability of

$$1 - (1 - p)^k \ge 1 - e^{-ck \ln n/n}$$

of being one. Note that $n - (|S_1| + |S_2| + \dots + |S_i|) \ge 999n/1000$. So,

$$E(|S_{i+1}|) \ge \frac{999n}{1000}(1 - e^{-ck\frac{\ln n}{n}}).$$

Subtracting 200k from each side

$$E(|S_{i+1}|) - 200k \ge \frac{n}{2} \left(1 - e^{-ck\frac{\ln n}{n}} - 400\frac{k}{n} \right).$$

Let $\alpha = \frac{k}{n}$ and $f(\alpha) = 1 - e^{-c\alpha \ln n} - 400\alpha$. By differentiation $f''(\alpha) \leq 0$, so f is concave and the minimum value of f over the interval [0, 1/1000] is attained at one of the end points. It is easy to check that both f(0) and f(1/1000) are greater than or equal to zero for sufficiently large n. Thus, f is nonnegative throughout the interval proving that $E(|S_{i+1}|) \geq 200|S_i|$. The lemma follows from Chernoff bounds.

Theorem 4.17 For $p \ge c \ln n/n$, where c is a sufficiently large constant, almost surely, G(n, p) has diameter $O(\ln n)$.

Proof: By Corollary 4.2, almost surely, the degree of every vertex is $\Omega(np) = \Omega(\ln n)$, which is at least $20 \ln n$ for c sufficiently large. Assume this holds. So, for a fixed vertex v, S_1 as defined in Lemma 4.16 satisfies $|S_1| \ge 20 \ln n$.

Let i_0 be the least i such that $|S_1|+|S_2|+\cdots+|S_i| > n/1000$. From Lemma 4.16 and the union bound, the probability that for some $i, 1 \le i \le i_0-1, |S_{i+1}| < 2(|S_1|+|S_2|+\cdots+|S_i|)$ is at most $\sum_{k=20 \ln n}^{n/1000} e^{-10k} \le 1/n^4$. So, with probability at least $1 - (1/n^4)$, each S_{i+1} is at least double the sum of the previous S_j 's, which implies that in $O(\ln n)$ steps, $i_0 + 1$ is reached.

Consider any other vertex w. We wish to find a short $O(\ln n)$ length path between v and w. By the same argument as above, the number of vertices at distance $O(\ln n)$ from w is at least n/1000. To complete the argument, either these two sets intersect in which case we have found a path from v to w of length $O(\ln n)$ or they do not intersect. In the latter case, with high probability there is some edge between them. For a pair of disjoint sets of size at least n/1000, the probability that none of the possible $n^2/10^6$ or more edges between them is present is at most $(1-p)^{n^2/10^6} = e^{-\Omega(n \ln n)}$. There are at most 2^{2n} pairs of such sets and so the probability that there is some such pair with no edges is $e^{-\Omega(n \ln n) + O(n)} \to 0$. Note that there is no conditioning problem since we are arguing this for every pair of such sets. Think of whether such an argument made for just the n subsets of vertices, which are vertices at distance at most $O(\ln n)$ from a specific vertex, would work.

4.6 Phase Transitions for Increasing Properties

For many graph properties such as connectivity, having no isolated vertices, having a cycle, etc., the probability of a graph having the property increases as edges are added to the graph. Such a property is called an increasing property. Q is an *increasing property* of graphs if when a graph G has the property, any graph obtained by adding edges to G must also have the property. In this section we show that any increasing property, in fact, has a threshold, although not necessarily a sharp one.

The notion of increasing property is defined in terms of adding edges. The following intuitive lemma proves that if Q is an increasing property, then increasing p in G(n, p) increases the probability of the property Q.

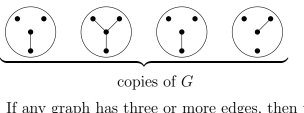
Lemma 4.18 If Q is an increasing property of graphs and $0 \le p \le q \le 1$, then the probability that G(n,q) has property Q is greater than or equal to the probability that G(n,p) has property Q.

Proof: This proof uses an interesting relationship between G(n, p) and G(n, q). Generate G(n,q) as follows. First generate G(n,p). This means generating a graph on n vertices with edge probabilities p. Then, independently generate another graph $G\left(n, \frac{q-p}{1-p}\right)$ and take the union by putting in an edge if either of the two graphs has the edge. Call the resulting graph H. The graph H has the same distribution as G(n,q). This follows since the probability that an edge is in H is $p + (1-p)\frac{q-p}{1-p} = q$, and, clearly, the edges of H are independent. The lemma follows since whenever G(n, p) has the property Q.

We now introduce a notion called *replication*. An *m*-fold replication of G(n, p) is a random graph obtained as follows. Generate *m* independent copies of G(n, p) on the same set of vertices. Include an edge in the *m*-fold replication if the edge is in any one of the *m* copies of G(n, p). The resulting random graph has the same distribution as G(n,q) where $q = 1 - (1-p)^m$ since the probability that a particular edge is not in the *m*-fold replication is the product of probabilities that it is not in any of the *m* copies of G(n, p). If the *m*-fold replication of G(n, p) does not have an increasing property Q, then none of the *m* copies of G(n, p) has the property. The converse is not true. If no copy has the property, their union may have it. Since Q is an increasing property and $q = 1 - (1-p)^m \leq 1 - (1-mp) = mp$

$$\operatorname{Prob}\left(G(n,mp) \operatorname{has} Q\right) \ge \operatorname{Prob}\left(G(n,q) \operatorname{has} Q\right) \tag{4.3}$$

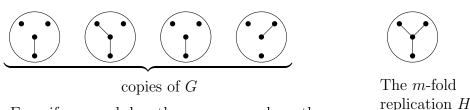
We now show that every increasing property Q has a phase transition. The transition occurs at the point at which the probability that G(n, p) has property Q is $\frac{1}{2}$. We will prove that for any function asymptotically less then p(n) that the probability of having property Q goes to zero as n goes to infinity.





The m-fold replication H

If any graph has three or more edges, then the m-fold replication has three or more edges.



Even if no graph has three or more edges, the m-fold replication might have three or more edges.

Figure 4.12: The property that G has three or more edges is an increasing property. Let H be the m-fold replication of G. If any copy of G has three or more edges, H has three or more edges. However, H can have three or more edges even if no copy of G has three or more edges.

Theorem 4.19 Every increasing property Q of G(n, p) has a phase transition at p(n), where for each n, p(n) is the minimum real number a_n for which the probability that $G(n, a_n)$ has property Q is 1/2.

Proof: Let $p_0(n)$ be any function such that

$$\lim_{n \to \infty} \frac{p_0(n)}{p(n)} = 0.$$

We assert that almost surely $G(n, p_0)$ does not have the property Q. Suppose for contradiction, that this is not true. That is, the probability that $G(n, p_0)$ has the property Q does not converge to zero. By the definition of a limit, there exists $\varepsilon > 0$ for which the probability that $G(n, p_0)$ has property Q is at least ε on an infinite set I of n. Let $m = \lceil (1/\varepsilon) \rceil$. Let G(n, q) be the *m*-fold replication of $G(n, p_0)$. The probability that G(n, q) does not have Q is at most $(1 - \varepsilon)^m \leq e^{-1} \leq 1/2$ for all $n \in I$. For these n, by (11.4)

 $\operatorname{Prob}(G(n, mp_0) \text{ has } Q) \ge \operatorname{Prob}(G(n, q) \text{ has } Q) \ge 1/2.$

Since p(n) is the minimum real number a_n for which the probability that $G(n, a_n)$ has property Q is 1/2, it must be that $mp_0(n) \ge p(n)$. This implies that $\frac{p_0(n)}{p(n)}$ is at least 1/m infinitely often, contradicting the hypothesis that $\lim_{n\to\infty} \frac{p_0(n)}{p(n)} = 0$.

A symmetric argument shows that for any $p_1(n)$ such that $\lim_{n\to\infty} \frac{p(n)}{p_1(n)} = 0$, $G(n, p_1)$ almost surely has property Q.

4.7 Phase Transitions for CNF-sat

Phase transitions occur not only in random graphs, but in other random structures as well. An important example is that of satisfiability for a Boolean formula in conjunctive normal form.

Generate a random CNF formula f with n variables, m clauses, and k literals per clause, where a literal is a variable or its negation. Specifically, each clause in f is selected independently at random from the set of all $\binom{n}{k}2^k$ possible clauses of size k. Equivalently, to generate a clause, choose a random set of k distinct variables, and then for each of those variables choose to either negate it or not with equal probability. Here, the number of clauses n is going to infinity, m is a function of n, and k is a fixed constant. A reasonable value to think of for k is k = 3. Unsatisfiability is an increasing property since adding more clauses preserves unsatisfiability. By arguments similar to the last section, there is a phase transition, i.e., a function m(n) such that if $m_1(n)$ is o(m(n)), a random formula with $m_1(n)$ clauses is, almost surely, satisfiable and for $m_2(n)$ with $m_2(n)/m(n) \to \infty$, a random formula with $m_2(n)$ clauses is, almost surely, unsatisfiable. It has been conjectured that there is a constant r_k independent of n such that $r_k n$ is a sharp threshold.

Here we derive upper and lower bounds on r_k . It is relatively easy to get an upper bound on r_k . A fixed truth assignment satisfies a random k clause with probability $1 - \frac{1}{2^k}$ because of the 2^k truth assignments to the k variables in the clause, only one fails to satisfy the clause. Thus, with probability $\frac{1}{2^k}$, the clause is not satisfied, and with probability $1 - \frac{1}{2^k}$, the clause is satisfied. Let m = cn. Now, cn independent clauses are all satisfied by the fixed assignment with probability $(1 - \frac{1}{2^k})^{cn}$. Since there are 2^n truth assignments, the expected number of satisfying assignments for a formula with cn clauses is $2^n (1 - \frac{1}{2^k})^{cn}$. If $c = 2^k \ln 2$, the expected number of satisfying assignments is

$$2^n \left(1 - \frac{1}{2^k}\right)^{n2^k \ln 2}$$

 $\left(1-\frac{1}{2^k}\right)^{2^k}$ is at most 1/e and approaches 1/e in the limit. Thus,

$$2^n \left(1 - \frac{1}{2^k}\right)^{n2^k \ln 2} \le 2^n e^{-n \ln 2} = 2^n 2^{-n} = 1.$$

For $c > 2^k \ln 2$, the expected number of satisfying assignments goes to zero as $n \to \infty$. Here the expectation is over the choice of clauses which is random, not the choice of a truth assignment. From the first moment method, it follows that a random formula with cn clauses is almost surely not satisfiable. Thus, $r_k \leq 2^k \ln 2$.

The other direction, showing a lower bound for r_k , is not that easy. From now on, we focus only on the case k = 3. The statements and algorithms given here can be extended

to $k \ge 4$, but with different constants. It turns out that the second moment method cannot be directly applied to get a lower bound on r_3 because the variance is too high. A simple algorithm, called the Smallest Clause Heuristic (abbreviated SC), yields a satisfying assignment with probability tending to one if $c < \frac{2}{3}$, proving that $r_3 \ge \frac{2}{3}$. Other more difficult to analyze algorithms, push the lower bound on r_3 higher.

The Smallest Clause Heuristic repeatedly executes the following. Assign true to a random literal in a random smallest length clause and delete the clause since it is now satisfied. In more detail, pick at random a 1-literal clause, if one exists, and set that literal to true. If there is no 1-literal clause, pick a 2-literal clause, select one of its two literals and set the literal to true. Otherwise, pick a 3-literal clause and a literal in it and set the literal to true. If we encounter a 0-length clause, then we have failed to find a satisfying assignment; otherwise, we have found one.

A related heuristic, called the Unit Clause Heuristic, selects a random clause with one literal, if there is one, and sets the literal in it to true. Otherwise, it picks a random as yet unset literal and sets it to true. Another variation is the "pure literal" heuristic. It sets a random "pure literal", a literal whose negation does not occur in any clause, to true, if there are any pure literals; otherwise, it sets a random literal to true.

When a literal w is set to true, all clauses containing w are deleted, since they are satisfied, and \bar{w} is deleted from any clause containing \bar{w} . If a clause is reduced to length zero (no literals), then the algorithm has failed to find a satisfying assignment to the formula. The formula may, in fact, be satisfiable, but the algorithm has failed.

Example: Consider a 3-CNF formula with n variables and cn clauses. With n variables there are 2n literals, since a variable and its complement are distinct literals. The expected number of times a literal occurs is calculated as follows. Each clause has three literals. Thus, each of the 2n different literals occurs $\frac{(3cn)}{2n} = \frac{3}{2}c$ times on average. Suppose c = 5. Then each literal appears 7.5 times on average. If one sets a literal to true, one would expect to satisfy 7.5 clauses. However, this process is not repeatable since after setting a literal to true there is conditioning so that the formula is no longer random.

Theorem 4.20 If the number of clauses in a random 3-CNF formula grows as cn where c is a constant less than 2/3, then with probability 1 - o(1), the Shortest Clause (SC) Heuristic finds a satisfying assignment.

The proof of this theorem will take the rest of the section. A general impediment to proving that simple algorithms work for random instances of many problems is conditioning. At the start, the input is random and has properties enjoyed by random instances. But, as the algorithm is executed, the data is no longer random; it is conditioned on the steps of the algorithm so far. In the case of SC and other heuristics for finding a satisfying assignment for a Boolean formula, the argument to deal with conditioning is relatively simple.

We supply some intuition before going to the proof. Imagine maintaining a queue of 1 and 2-clauses. A 3-clause enters the queue when one of its literals is set to false and it becomes a 2-clause. SC always picks a 1 or 2-clause if there is one and sets one of its literals to true. At any step when the total number of 1 and 2-clauses is positive, one of the clauses is removed from the queue. Consider the arrival rate, that is, the expected number of arrivals into the queue at a given time t. For a particular clause to arrive into the queue at time t to become a 2-clause, it must contain the negation of the literal being set to true at time t. It can contain any two other literals not yet set. The number of such clauses is $\binom{n-t}{2}2^2$. So, the probability that a particular clause arrives in the queue at time t is at most

$$\frac{\binom{n-t}{2}2^2}{\binom{n}{3}2^3} \le \frac{3}{2(n-2)}.$$

Since there are cn clauses in total, the arrival rate is $\frac{3c}{2}$, which for c < 2/3 is a constant strictly less than one. The arrivals into the queue of different clauses occur independently (Lemma 4.21), the queue has arrival rate strictly less than one, and the queue loses one or more clauses whenever it is nonempty. This implies that the queue never has too many clauses in it. A slightly more complicated argument will show that no clause remains as a 1 or 2-clause for $\omega(\ln n)$ steps (Lemma 4.22). This implies that the probability of two contradictory 1-length clauses, which is a precursor to a 0-length clause, is very small.

Lemma 4.21 Let T_i be the first time that clause *i* turns into a 2-clause. T_i is ∞ if clause *i* gets satisfied before turning into a 2-clause. The T_i are mutually independent over the randomness in constructing the formula and the randomness in SC, and for any *t*,

$$Prob(T_i = t) \le \frac{3}{2(n-2)}.$$

Proof: For the proof, generate the clauses in a different way. The important thing is that the new method of generation, called the method of "deferred decisions", results in the same distribution of input formulae as the original. The method of deferred decisions is tied in with the SC algorithm and works as follows. At any time, the length of each clause (number of literals) is all that we know; we have not yet picked which literals are in each clause. At the start, every clause has length three and SC picks one of the clauses uniformly at random. Now, SC wants to pick one of the three literals in that clause to set to true, but we do not know which literals are in the clause. At this point, we pick uniformly at random one of the 2n possible literals. Say for illustration, we picked \bar{x}_{102} . The literal \bar{x}_{102} is placed in the clause and set to true. The literal x_{102} is set to false. We must also deal with occurrences of the literal or its negation in all other clauses, but again, we do not know which clauses have such an occurrence. We decide that now. For each clause, independently, with probability 3/n include either the literal \bar{x}_{102} or its negation x_{102} , each with probability 1/2. In the case that we included \bar{x}_{102} (the literal we had set to false),

we decrease the residual length of the clause by one.

At a general stage, suppose the fates of i variables have already been decided and n-i remain. The residual length of each clause is known. Among the clauses that are not yet satisfied, choose a random shortest length clause. Among the n-i variables remaining, pick one uniformly at random, then pick it or its negation as the new literal. Include this literal in the clause thereby satisfying it. Since the clause is satisfied, the algorithm deletes it. For each other clause, do the following. If its residual length is l, decide with probability l/(n-i) to include the new variable in the clause and if so with probability 1/2 each, include it or its negation. If literal that was set to true is included in a clause, delete the clause as it is now satisfied. If its negation is included in a clause, then just delete the literal and decrease the residual length of the clause by one.

Why does this yield the same distribution as the original one? First, observe that the order in which the variables are picked by the method of deferred decisions is independent of the clauses; it is just a random permutation of the *n* variables. Look at any one clause. For a clause, we decide in order whether each variable or its negation is in the clause. So for a particular clause and a particular triple i, j, and k with i < j < k, the probability that the clause contains the i^{th} , the j^{th} , and k^{th} literal (or their negations) in the order determined by deferred decisions is:

$$\left(1 - \frac{3}{n}\right) \left(1 - \frac{3}{n-1}\right) \cdots \left(1 - \frac{3}{n-i+2}\right) \frac{3}{n-i+1} \left(1 - \frac{2}{n-i}\right) \left(1 - \frac{2}{n-i-1}\right) \cdots \left(1 - \frac{2}{n-j+2}\right) \frac{2}{n-j+1} \left(1 - \frac{1}{n-j}\right) \left(1 - \frac{1}{n-j-1}\right) \cdots \left(1 - \frac{1}{n-k+2}\right) \frac{1}{n-k+1} = \frac{3}{n(n-1)(n-2)},$$

where the $(1 - \cdots)$ factors are for not picking the current variable or negation to be included and the others are for including the current variable or its negation. Independence among clauses follows from the fact that we have never let the occurrence or nonoccurrence of any variable in any clause influence our decisions on other clauses.

Now, we prove the lemma by appealing to the method of deferred decisions to generate the formula. $T_i = t$ if and only if the method of deferred decisions does not put the current literal at steps $1, 2, \ldots, t - 1$ into the i^{th} clause, but puts the negation of the literal at step t into it. Thus, the probability is precisely

$$\frac{1}{2}\left(1-\frac{3}{n}\right)\left(1-\frac{3}{n-1}\right)\cdots\left(1-\frac{3}{n-t+2}\right)\frac{3}{n-t+1} \le \frac{3}{2(n-2)},$$

as claimed. Clearly the T_i are independent since again deferred decisions deal with different clauses independently.

Lemma 4.22 There exists a constant c_2 such that with probability 1 - o(1), no clause remains a 2 or 1-clause for more than $c_2 \ln n$ steps. I.e., once a 3-clause becomes a 2-clause, it is either satisfied or reduced to a 0-clause in $O(\ln n)$ steps. **Proof:** Say that t is a "busy time" if there exists at least one 2-clause or 1-clause at time t, and define a time-window [r + 1, s] to be a "busy window" if time r is not busy but then each $t \in [r + 1, s]$ is a busy time. We will prove that for some constant c_2 , with probability 1 - o(1), all busy windows have length at most $c_2 \ln n$.

Fix some r, s and consider the event that [r + 1, s] is a busy window. Since SC always decreases the total number of 1 and 2-clauses by one whenever it is positive, we must have generated at least s - r new 2-clauses between r and s. Now, define an indicator random variable for each 3-clause which has value one if the clause turns into a 2-clause between r and s. By Lemma 4.21 these variables are independent and the probability that a particular 3-clause turns into a 2-clause at a time t is at most 3/(2(n-2)). Summing over t between r and s,

Prob (a 3-clause turns into a 2-clause during
$$[r, s]$$
) $\leq \frac{3(s-r)}{2(n-2)}$.

Since there are cn clauses in all, the expected sum of the indicator random variables is $cn\frac{3(s-r)}{2(n-2)} \approx \frac{3c(s-r)}{2}$. Note that 3c/2 < 1, which implies the arrival rate into the queue of 2 and 1-clauses is a constant strictly less than one. Using Chernoff bounds, if $s-r \ge c_2 \ln n$ for appropriate constant c_2 , the probability that more than s-r clauses turn into 2-clauses between r and s is at most $1/n^3$. Applying the union bound over all $O(n^2)$ possible choices of r and s, we get that the probability that any clause remains a 2 or 1-clause for more than $c_2 \ln n$ steps is o(1).

Now, assume the 1 - o(1) probability event of Lemma 4.22 that no clause remains a 2 or 1-clause for more than $c_2 \ln n$ steps. We will show that this implies it is unlikely the SC algorithm terminates in failure.

Suppose SC terminates in failure. This means that at some time t, the algorithm generates a 0-clause. At time t-1, this clause must have been a 1-clause. Suppose the clause consists of the literal w. Since at time t-1, there is at least one 1-clause, the shortest clause rule of SC selects a 1-clause and sets the literal in that clause to true. This other clause must have been \bar{w} . Let t_1 be the first time either of these two clauses, w or \bar{w} , became a 2-clause. We have $t - t_1 \leq c_2 \ln n$. Clearly, until time t, neither of these two clauses is picked by SC. So, the literals which are set to true during this period are chosen independent of these clauses. Say the two clauses were w + x + y and $\bar{w} + u + v$ at the start. x, y, u, and v must all be negations of literals set to true during steps t_1 to t. So, there are only $O((\ln n)^4)$ choices for x, y, u, and v for a given value of t. There are O(n) choices of w, $O(n^2)$ choices of which two clauses i, j of the input become these w and \bar{w} , and n choices for t. Thus, there are $O(n^4(\ln n)^4)$ choices for what these clauses contain and which clauses they are in the input. On the other hand, for any given i, j, the probability that clauses i, j both match a given set of literals is $O(1/n^6)$. Thus the probability that these choices are actually realized is therefore $O\left(n^4(\ln n)^4/n^6\right) = o(1)$, as required.

4.8 Nonuniform and Growth Models of Random Graphs

4.8.1 Nonuniform Models

So far we have considered the random graph G(n, p) in which all vertices have the same expected degree and showed that the degree is concentrated close to its expectation. However, large graphs occurring in the real world tend to have power law degree distributions. For a power law degree distribution, the number f(d) of vertices of degree d plotted as a function of d satisfies $f(d) \leq c/d^{\alpha}$, where α and c are constants.

To generate such graphs, we stipulate that there are f(d) vertices of degree d and choose uniformly at random from the set of graphs with this degree distribution. Clearly, in this model the graph edges are not independent and this makes these random graphs harder to analyze. But the question of when phase transitions occur in random graphs with arbitrary degree distributions is still of interest. In this section, we consider when a random graph with a nonuniform degree distribution has a giant component. Our treatment in this section, and subsequent ones, will be more intuitive without providing rigorous proofs.

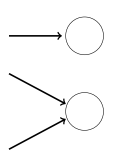
4.8.2 Giant Component in Random Graphs with Given Degree Distribution

Molloy and Reed address the issue of when a random graph with a nonuniform degree distribution has a giant component. Let λ_i be the fraction of vertices of degree *i*. There will be a giant component if and only if $\sum_{i=0}^{\infty} i(i-2)\lambda_i > 0$.

To see intuitively that this is the correct formula, consider exploring a component of a graph starting from a given seed vertex. Degree zero vertices do not occur except in the case where the vertex is the seed. If a degree one vertex is encountered, then that terminates the expansion along the edge into the vertex. Thus, we do not want to encounter too many degree one vertices. A degree two vertex is neutral in that the vertex is entered by one edge and left by the other. There is no net increase in the size of the frontier. Vertices of degree *i* greater than two increase the frontier by i - 2 vertices. The vertex is entered by one of its edges and thus there are i - 1 edges to new vertices in the frontier for a net gain of i - 2. The $i\lambda_i$ in $i(i - 2)\lambda_i$ is proportional to the probability of reaching a degree *i* vertex and the i - 2 accounts for the increase or decrease in size of the frontier when a degree *i* vertex is reached.

Example: Consider applying the Molloy Reed conditions to the G(n, p) model, and use p_i to denote the probability that a vertex has degree i, i.e., to analog to λ_i . It turns out that the summation $\sum_{i=0}^{n} i(i-2)p_i$ gives value zero precisely when p = 1/n, the point at which the phase transition occurs. At p = 1/n, the average degree of each vertex is one and there are n/2 edges. However, the actual degree distribution of the vertices is binomial, where the probability that a vertex is of degree i is given by $p_i = {n \choose i} p^i (1-p)^{n-i}$.

We now show that $\lim_{n \to \infty} \sum_{i=0}^{n} i(i-2)p_i = 0$ for $p_i = \binom{n}{i}p^i(1-p)^{n-i}$ when p = 1/n.



Consider a graph in which half of the vertices are degree one and half are degree two. If a vertex is selected at random, it is equally likely to be degree one or degree two. However, if we select an edge at random and walk to a random endpoint, the vertex is twice as likely to be degree two as degree one. In many graph algorithms, a vertex is reached by randomly selecting an edge and traversing the edge to reach an endpoint. In this case, the probability of reaching a degree *i* vertex is proportional to $i\lambda_i$ where λ_i is the fraction of vertices that are degree *i*.

Figure 4.13: Probability of encountering a degree d vertex when following a path in a graph.

$$\lim_{n \to \infty} \sum_{i=0}^{n} i(i-2) \binom{n}{i} \left(\frac{1}{n}\right)^{i} \left(1-\frac{1}{n}\right)^{n-i}$$

$$= \lim_{n \to \infty} \sum_{i=0}^{n} i(i-2) \frac{n(n-1)\cdots(n-i+1)}{i! n^{i}} \left(1-\frac{1}{n}\right)^{n} \left(1-\frac{1}{n}\right)^{-i}$$

$$= \frac{1}{e} \lim_{n \to \infty} \sum_{i=0}^{n} i(i-2) \frac{n(n-1)\cdots(n-i+1)}{i! n^{i}} \left(\frac{n}{n-1}\right)^{i}$$

$$\leq \sum_{i=0}^{\infty} \frac{i(i-2)}{i!}.$$

To see that $\sum_{i=0}^{\infty} \frac{i(i-2)}{i!} = 0$, note that

$$\sum_{i=0}^{\infty} \frac{i}{i!} = \sum_{i=1}^{\infty} \frac{i}{i!} = \sum_{i=1}^{\infty} \frac{1}{(i-1)!} = \sum_{i=0}^{\infty} \frac{1}{i!}$$

and

$$\sum_{i=0}^{\infty} \frac{i^2}{i!} = \sum_{i=1}^{\infty} \frac{i}{(i-1)!} = \sum_{i=0}^{\infty} \frac{i+1}{i!} = \sum_{i=0}^{\infty} \frac{i}{i!} + \sum_{i=0}^{\infty} \frac{1}{i!} = 2\sum_{i=0}^{\infty} \frac{1}{i!}.$$

Thus,

$$\sum_{i=0}^{\infty} \frac{i(i-2)}{i!} = \sum_{i=0}^{\infty} \frac{i^2}{i!} - 2\sum_{i=0}^{\infty} \frac{i}{i!} = 0.$$

4.9 Growth Models

4.9.1 Growth Model Without Preferential Attachment

Many graphs that arise in the outside world started as small graphs that grew over time. In a model for such graphs, vertices and edges are added to the graph over time. In such a model there are many ways in which to select the vertices for attaching a new edge. One is to select two vertices uniformly at random from the set of existing vertices. Another is to select two vertices with probability proportional to their degree. This latter method is referred to as preferential attachment. A variant of this method would be to add a new vertex at each unit of time and with probability δ add an edge where one end of the edge is the new vertex and the other end is a vertex selected with probability proportional to its degree. The graph generated by this latter method is a tree.

Consider a growth model for a random graph without preferential attachment. Start with zero vertices at time zero. At each unit of time a new vertex is created and with probability δ , two vertices chosen at random are joined by an edge. The two vertices may already have an edge between them. In this case, we will add another edge. So, the resulting structure is a multi-graph, rather then a graph. But since at time t, there are t vertices and in expectation only $O(\delta t)$ edges where there are t^2 pairs of vertices, it is very unlikely that there will be many multiple edges.

The degree distribution for this growth model is calculated as follows. The number of vertices of degree k at time t is a random variable. Let $d_k(t)$ be the expectation of the number of vertices of degree k at time t. The number of isolated vertices increases by one at each unit of time and decreases by the number of isolated vertices, b(t), that are picked to be end points of the new edge. b(t) can take on values 0,1, or 2. Taking expectations,

$$d_0(t+1) = d_0(t) + 1 - E(b(t)).$$

Now b(t) is the sum of two 0-1 valued random variables whose values are the number of degree zero vertices picked for each end point of the new edge. Even though the two random variables are not independent, the expectation of b(t) is the sum of the expectations of the two variables and is $2\delta \frac{d_0(t)}{t}$. Thus,

$$d_0(t+1) = d_0(t) + 1 - 2\delta \frac{d_0(t)}{t}.$$

The number of degree k vertices increases whenever a new edge is added to a degree k-1 vertex and decreases when a new edge is added to a degree k vertex. Reasoning as above,

$$d_k(t+1) = d_k(t) + 2\delta \frac{d_{k-1}(t)}{t} - 2\delta \frac{d_k(t)}{t}.$$
(4.4)

Note that this formula, as others in this section, is not quite precise. For example, the same vertex may be picked twice, so that the new edge is a self-loop. For $k \ll t$, this

problem contributes a minuscule error. Restricting k to be a fixed constant and letting $t \to \infty$ in this section avoids these problems.

Assume that the above equations are exactly valid. Clearly, $d_0(1) = 1$ and $d_1(1) = d_2(1) = \cdots = 0$. By induction on t, there is a unique solution to (4.4), since given $d_k(t)$ for all k, the equation determines $d_k(t+1)$ for all k. There is a solution of the form $d_k(t) = p_k t$, where p_k depends only on k and not on t, provided k is fixed and $t \to \infty$. Again, this is not precisely true since $d_1(1) = 0$ and $d_1(2) > 0$ clearly contradict the existence of a solution of the form $d_1(t) = p_1 t$.

Set $d_k(t) = p_k t$. Then,

$$(t+1) p_0 = p_0 t + 1 - 2\delta \frac{p_0 t}{t}$$
$$p_0 = 1 - 2\delta p_0$$
$$p_0 = \frac{1}{1+2\delta}$$

and

$$(t+1) p_k = p_k t + 2\delta \frac{p_{k-1}t}{t} - 2\delta \frac{p_k t}{t}$$
$$p_k = 2\delta p_{k-1} - 2\delta p_k$$

$$p_{k} = \frac{2\delta}{1+2\delta} p_{k-1}$$

$$= \left(\frac{2\delta}{1+2\delta}\right)^{k} p_{0}$$

$$= \frac{1}{1+2\delta} \left(\frac{2\delta}{1+2\delta}\right)^{k}.$$
(4.5)

Thus, the model gives rise to a graph with a degree distribution that falls off exponentially fast with degree.

The generating function for component size

Let $n_k(t)$ be the expected number of components of size k at time t. Then $n_k(t)$ is proportional to the probability that a randomly picked component is of size k. This is not the same as picking the component containing a randomly selected vertex (see Figure 4.14). Indeed, the probability that the size of the component containing a randomly selected vertex is k is proportional to $kn_k(t)$. We will show that there is a solution for $n_k(t)$ of the form $a_k t$ where a_k is a constant independent of t. After showing this, we focus on the generating function g(x) for the numbers $ka_k(t)$ and use g(x) to find the threshold for giant components.

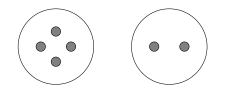


Figure 4.14: In selecting a component at random, each of the two components is equally likely to be selected. In selecting the component containing a random vertex, the larger component is twice as likely to be selected.

Consider $n_1(t)$, the expected number of isolated vertices at time t. At each unit of time, an isolated vertex is added to the graph and an expected $\frac{2\delta n_1(t)}{t}$ many isolated vertices are chosen for attachment and thereby leave the set of isolated vertices. Thus,

$$n_1(t+1) = n_1(t) + 1 - 2\delta \frac{n_1(t)}{t}.$$

For k > 1, $n_k(t)$ increases when two smaller components whose sizes sum to k are joined by an edge and decreases when a vertex in a component of size k is chosen for attachment. The probability that a vertex selected at random will be in a size k component is $\frac{kn_k(t)}{t}$. Thus,

$$n_k(t+1) = n_k(t) + \delta \sum_{j=1}^{k-1} \frac{jn_j(t)}{t} \frac{(k-j)n_{k-j}(t)}{t} - 2\delta \frac{kn_k(t)}{t}.$$

To be precise, one needs to consider the actual number of components of various sizes, rather than the expected numbers. Also, if both vertices at the end of the edge are in the same k-vertex component, then $n_k(t)$ does not go down as claimed. These small inaccuracies can be ignored.

Consider solutions of the form $n_k(t) = a_k t$. Note that $n_k(t) = a_k t$ implies the number of vertices in a connected component of size k is $ka_k t$. Since the total number of vertices at time t is t, ka_k is the probability that a random vertex is in a connected component of size k. The recurrences here are valid only for k fixed as $t \to \infty$. So $\sum_{k=0}^{\infty} ka_k$ may be less than 1, in which case, there are nonfinite size components whose sizes are growing with t. Solving for a_k yields $a_1 = \frac{1}{1+2\delta}$ and $a_k = \frac{\delta}{1+2k\delta} \sum_{j=1}^{k-1} j(k-j)a_ja_{k-j}$.

Consider the generating function g(x) for the distribution of component sizes where the coefficient of x^k is the probability that a vertex chosen at random is in a component of size k.

$$g(x) = \sum_{k=1}^{\infty} k a_k x^k.$$

Now, $g(1) = \sum_{k=0}^{\infty} ka_k$ is the probability that a randomly chosen vertex is in a finite sized component. For $\delta = 0$, this is clearly one, since all vertices are in components of size

one. On the other hand, for $\delta = 1$, the vertex created at time one has expected degree $\log n$ (since its expected degree increases by 2/t and $\sum_{t=1}^{n} (2/t) = \Theta(\log n)$); so, it is in a nonfinite size component. This implies that for $\delta = 1$, g(1) < 1 and there is a nonfinite size component. Assuming continuity, there is a $\delta_{critical}$ above which g(1) < 1. From the formula for the $a'_i s$, we will derive the differential equation

$$g = -2\delta xg' + 2\delta xgg' + x$$

and then use the equation for g to determine the value of δ at which the phase transition for the appearance of a nonfinite sized component occurs.

Derivation of g(x)

From

$$a_1 = \frac{1}{1+2\delta}$$

and

$$a_k = \frac{\delta}{1+2k\delta} \sum_{j=1}^{k-1} j(k-j)a_j a_{k-j}$$

derive the equations

$$a_1 \left(1 + 2\delta \right) - 1 = 0$$

and

$$a_k \left(1 + 2k\delta\right) = \delta \sum_{j=1}^{k-1} j(k-j)a_j a_{k-j}$$

for $k \geq 2$. The generating function is formed by multiplying the k^{th} equation by kx^k and summing over all k. This gives

$$-x + \sum_{k=1}^{\infty} ka_k x^k + 2\delta x \sum_{k=1}^{\infty} a_k k^2 x^{k-1} = \delta \sum_{k=1}^{\infty} kx^k \sum_{j=1}^{k-1} j(k-j)a_j a_{k-j}$$

Note that

$$g(x) = \sum_{k=1}^{\infty} k a_k x^k$$
 and $g'(x) = \sum_{k=1}^{\infty} a_k k^2 x^{k-1}$.

Thus,

$$-x + g(x) + 2\delta x g'(x) = \delta \sum_{k=1}^{\infty} k x^k \sum_{j=1}^{k-1} j(k-j) a_j a_{k-j}.$$

Working with the right hand side

$$\delta \sum_{k=1}^{\infty} kx^k \sum_{j=1}^{k-1} j(k-j)a_j a_{k-j} = \delta x \sum_{k=1}^{\infty} \sum_{j=1}^{k-1} j(k-j)(j+k-j)x^{k-1}a_j a_{k-j}.$$

Now breaking the j + k - j into two sums gives

$$\delta x \sum_{k=1}^{\infty} \sum_{j=1}^{k-1} j^2 a_j x^{j-1} (k-j) a_{k-j} x^{k-j} + \delta x \sum_{k=1}^{\infty} \sum_{j=1}^{k-1} j a_j x^j (k-j)^2 a_{k-j} x^{k-j-1}.$$

Notice that the second sum is obtained from the first by substituting k - j for j and that both terms are $\delta x g' g$. Thus,

$$-x + g(x) + 2\delta x g'(x) = 2\delta x g'(x)g(x).$$

Hence,

$$g' = \frac{1}{2\delta} \frac{1 - \frac{g}{x}}{1 - g}.$$

Phase transition for nonfinite components

The generating function g(x) contains information about the finite components of the graph. A finite component is a component of size $1, 2, \ldots$ which does not depend on t. Observe that $g(1) = \sum_{k=0}^{\infty} ka_k$ and hence g(1) is the probability that a randomly chosen vertex will belong to a component of finite size. If g(1) = 1 there are no nonfinite components. When $g(1) \neq 1$, then 1 - g(1) is the expected fraction of the vertices that are in nonfinite components. Potentially, there could be many such nonfinite components. But an argument similar to Part 3 of Theorem 4.8 concludes that two fairly large components would merge into one. Suppose there are two connected components at time t, each of size at least $t^{4/5}$. Consider the earliest created $\frac{1}{2}t^{4/5}$ vertices in each part. These vertices must have lived for at least $\frac{1}{2}t^{4/5}$ time after creation. At each time, the probability of an edge forming between two such vertices, one in each component, is at least $\delta\Omega(t^{-2/5})$ and so the probability that no such edge formed is at most $(1 - \delta t^{-2/5})^{t^{4/5/2}} \leq e^{-\Omega(\delta t^{2/5})} \to 0$. So with high probability of many components would have merged into one. But this still leaves open the possibility of many components of size t^{ε} , $(\ln t)^2$, or some other slowly growing function of t.

We now calculate the value of δ at which the phase transition for a nonfinite component occurs. Recall that the generating function for g(x) satisfies

$$g'(x) = \frac{1}{2\delta} \frac{1 - \frac{g(x)}{x}}{1 - g(x)}.$$

If δ is greater than some $\delta_{critical}$, then $g(1) \neq 1$. In this case the above formula at x = 1 simplifies with 1 - g(1) canceling from the numerator and denominator, leaving just $\frac{1}{2\delta}$. Since ka_k is the probability that a randomly chosen vertex is in a component of size k, the average size of the finite components is $g'(1) = \sum_{k=1}^{\infty} k^2 a_k$. Now, g'(1) is given by

$$g'(1) = \frac{1}{2\delta} \tag{4.6}$$

for all δ greater than $\delta_{critical}$. If δ is less than $\delta_{critical}$, then all vertices are in finite components. In this case g(1) = 1 and both the numerator and the denominator approach zero. Appling L'Hopital's rule

$$\lim_{x \to 1} g'(x) = \frac{1}{2\delta} \left. \frac{\frac{xg'(x) - g(x)}{x^2}}{g'(x)} \right|_{x=1}$$

or

$$(g'(1))^2 = \frac{1}{2\delta} (g'(1) - g(1)).$$

The quadratic $(g'(1))^2 - \frac{1}{2\delta}g'(1) + \frac{1}{2\delta}g(1) = 0$ has solutions

$$g'(1) = \frac{\frac{1}{2\delta} \pm \sqrt{\frac{1}{4\delta^2} - \frac{4}{2\delta}}}{2} = \frac{1 \pm \sqrt{1 - 8\delta}}{4\delta}.$$
(4.7)

The two solutions given by (4.7) become complex for $\delta > 1/8$ and thus can be valid only for $0 \le \delta \le 1/8$. For $\delta > 1/8$, the only solution is $g'(1) = \frac{1}{2\delta}$ and a nonfinite component exists. As δ is decreased, at $\delta = 1/8$ there is a singular point where for $\delta < 1/8$ there are three possible solutions, one from (4.6) which implies a giant component and two from (4.7) which imply no giant component. To determine which one of the three solutions is valid, consider the limit as $\delta \to 0$. In the limit all components are of size one since there are no edges. Only (4.7) with the minus sign gives the correct solution

$$g'(1) = \frac{1 - \sqrt{1 - 8\delta}}{4\delta} = \frac{1 - \left(1 - \frac{1}{2}8\delta - \frac{1}{4}64\delta^2 + \cdots\right)}{4\delta} = 1 + 4\delta + \cdots = 1.$$

In the absence of any nonanalytic behavior in the equation for g'(x) in the region $0 \le \delta < 1/8$, we conclude that (4.7) with the minus sign is the correct solution for $0 \le \delta < 1/8$ and hence the critical value of δ for the phase transition is 1/8. As we shall see, this is different from the static case.

As the value of δ is increased, the average size of the finite components increase from one to

$$\frac{1-\sqrt{1-8\delta}}{4\delta}\bigg|_{\delta=1/8} = 2$$

when δ reaches the critical value of 1/8. At $\delta = 1/8$, the average size of the finite components jumps to $\frac{1}{2\delta}\Big|_{\delta=1/8} = 4$ and then decreases as $\frac{1}{2\delta}$ as the giant component swallows up the finite components starting with the larger components.

Comparison to static random graph

Consider a static random graph with the same degree distribution as the graph in the growth model. Again let p_k be the probability of a vertex being of degree k. From (4.5)

$$p_k = \frac{(2\delta)^k}{(1+2\delta)^{k+1}} \,.$$

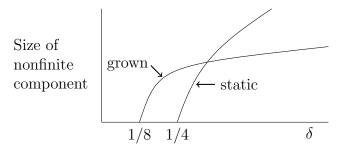


Figure 4.15: Comparison of the static random graph model and the growth model. The curve for the growth model is obtained by integrating g'.

Recall the Molloy Reed analysis of random graphs with given degree distributions which asserts that there is a phase transition at $\sum_{i=0}^{\infty} i(i-2)p_i = 0$. Using this, it is easy to see that a phase transition occurs for $\delta = 1/4$. For $\delta = 1/4$,

$$p_k = \frac{(2\delta)^k}{(1+2\delta)^{k+1}} = \frac{\left(\frac{1}{2}\right)^k}{\left(1+\frac{1}{2}\right)^{k+1}} = \frac{\left(\frac{1}{2}\right)^k}{\frac{3}{2}\left(\frac{3}{2}\right)^k} = \frac{2}{3}\left(\frac{1}{3}\right)^k$$

and

$$\sum_{i=0}^{\infty} i(i-2)\frac{2}{3} \left(\frac{1}{3}\right)^i = \frac{2}{3} \sum_{i=0}^{\infty} i^2 \left(\frac{1}{3}\right)^i - \frac{4}{3} \sum_{i=0}^{\infty} i \left(\frac{1}{3}\right)^i = \frac{2}{3} \times \frac{3}{2} - \frac{4}{3} \times \frac{3}{4} = 0.$$

Recall that $1 + a + a^2 + \dots = \frac{1}{1-a}$, $a + 2a^2 + 3a^3 \dots = \frac{a}{(1-a)^2}$, and $a + 4a^2 + 9a^3 \dots = \frac{a(1+a)}{(1-a)^3}$.

See references at end of the chapter for calculating the fractional size S_{static} of the giant component in the static graph. The result is

$$S_{static} = \begin{cases} 0 & \delta \le \frac{1}{4} \\ 1 - \frac{1}{\delta + \sqrt{\delta^2 + 2\delta}} & \delta > \frac{1}{4} \end{cases}$$

4.9.2 Growth Model With Preferential Attachment

Consider a growth model with preferential attachment. At each time unit, a vertex is added to the graph. Then with probability δ , an edge is attached to the new vertex and to a vertex selected at random with probability proportional to its degree. This model generates a tree with a power law distribution.

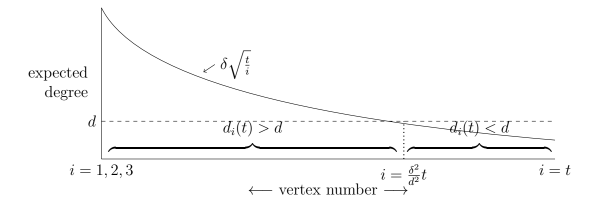


Figure 4.16: Illustration of degree of i^{th} vertex at time t. At time t, vertices numbered 1 to $\frac{\delta^2}{d^2}t$ have degrees greater than d.

Let $d_i(t)$ be the expected degree of the i^{th} vertex at time t. The sum of the expected degrees of all vertices at time t is $2\delta t$ and thus the probability that an edge is connected to vertex i at time t is $\frac{d_i(t)}{2\delta t}$. The degree of vertex i is governed by the equation

$$\frac{\partial}{\partial t}d_{i}(t) = \delta \frac{d_{i}\left(t\right)}{2\delta t} = \frac{d_{i}(t)}{2t}$$

where δ is the probability that an edge is added at time t and $\frac{d_i(t)}{2\delta t}$ is the probability that the vertex i is selected for the end point of the edge.

The two in the denominator governs the solution which is of the form $at^{\frac{1}{2}}$. The value of a is determined by the initial condition $d_i(t) = \delta$ at t = i. Thus, $\delta = ai^{\frac{1}{2}}$ or $a = \delta i^{-\frac{1}{2}}$. Hence, $d_i(t) = \delta \sqrt{\frac{t}{i}}$.

Next, we determine the probability distribution of vertex degrees. Now, $d_i(t)$ is less than d provided $i > \frac{\delta^2}{d^2}t$. The fraction of the t vertices at time t for which $i > \frac{\delta^2}{d^2}t$ and thus that the degree is less than d is $1 - \frac{\delta^2}{d^2}$. Hence, the probability that a vertex has degree less than d is $1 - \frac{\delta^2}{d^2}$. The probability density p(d) satisfies

$$\int_0^d p(d) \partial d = \operatorname{Prob}(\operatorname{degree} < d) = 1 - \frac{\delta^2}{d^2}$$

and can be obtained from the derivative of Prob(degree < d).

$$p(d) = \frac{\partial}{\partial d} \left(1 - \frac{\delta^2}{d^2} \right) = 2 \frac{\delta^2}{d^3},$$

a power law distribution.

4.10 Small World Graphs

In the 1960's, Stanley Milgram carried out an experiment that indicated that most pairs of individuals in the United States were connected by a short sequence of acquaintances. Milgram would ask a source individual, say in Nebraska, to start a letter on its journey to a target individual in Massachusetts. The Nebraska individual would be given basic information about the target including his address and occupation and asked to send the letter to someone he knew on a first name basis, who was closer to the target individual, in order to transmit the letter to the target in as few steps as possible. Each person receiving the letter would be given the same instructions. In successful experiments, it would take on average five to six steps for a letter to reach its target. This research generated the phrase "six degrees of separation" along with substantial research in social science on the interconnections between people. Surprisingly, there was no work on how to find the short paths using only local information.

In many situations, phenomena are modeled by graphs whose edges can be partitioned into local and long distance. We adopt a simple model of a directed graph due to Kleinberg, having local and long distance edges. Consider a 2-dimensional $n \times n$ grid where each vertex is connected to its four adjacent vertices via bidirectional local edges. In addition to these local edges, there is one long distance edge out of each vertex. The probability that the long distance edge from vertex u terminates at $v, v \neq u$, is a function of the distance d(u, v) from u to v. Here distance is measured by the shortest path consisting only of local grid edges. The probability is proportional to $1/d^r(u, v)$ for some constant r. This gives a one parameter family of random graphs. For r equal zero, $1/d^0(u,v) = 1$ for all u and v and thus the end of the long distance edge at u is uniformly distributed over all vertices independent of distance. As r increases the expected length of the long distance edge decreases. As r approaches infinity, there are no long distance edges and thus no paths shorter than that of the lattice path. What is interesting is that for r less than two, there are always short paths, but no local algorithm to find them. A local algorithm is an algorithm that is only allowed to remember the source, the destination, and its current location and can query the graph to find the long-distance edge at the current location. Based on this information, it decides the next vertex on the path.

The difficulty is that for r < 2, the end points of the long distance edges tend to be uniformly distributed over the vertices of the grid. Although short paths exist, it is unlikely on a short path to encounter a long distance edge whose end point is close to the destination. When r equals two, there are short paths and the simple algorithm that always selects the edge that ends closest to the destination will find a short path. For rgreater than two, again there is no local algorithm to find a short path. Indeed, with high probability, there are no short paths at all.

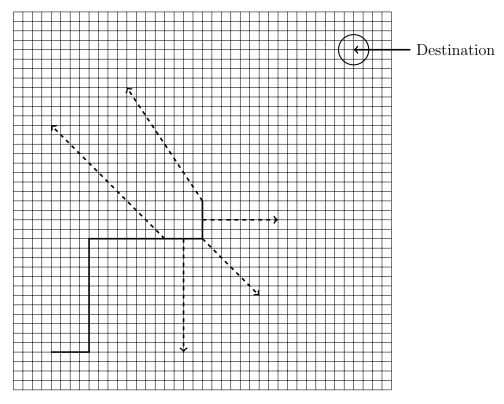


Figure 4.17: For d < 2, ends points of long distance edges are uniformly distributed and on a short path it is unlikely to encounter one that takes you close to the destination.

r>2 The lengths of long distance edges tend to be short so the probability of encountering a sufficiently long, long-distance edge is too low.

r = 2 Selecting the edge with end point closest to the destination finds a short path.

r < 2 The ends of long distance edges tend to be uniformly distributed. Short paths exist but a polylog length path is unlikely to encounter a long distance edge whose end point is close to the destination.

Figure 4.18: Effects of different values of r on the expected length of long distance edges and the ability to find short paths.

The probability that the long distance edge from u goes to v is proportional to $d^{-r}(u, v)$. Note that the constant of proportionality will vary with the vertex u depending on where u is relative to the border of the $n \times n$ grid. However, the number of vertices at distance exactly k from u is at most 4k and for $k \leq n/2$ is at least k. Let $c_r(u) = \sum_v d^{-r}(u, v)$ be the normalizing constant. It is the inverse of the constant of proportionality.

For r > 2, $c_r(u)$ is lower bounded by

$$c_r(u) = \sum_{v} d^{-r}(u, v) \ge \sum_{k=1}^{n/2} (k)k^{-r} = \sum_{k=1}^{n/2} k^{1-r} \ge 1.$$

No matter how large r is the first term of $\sum_{k=1}^{n/2} k^{1-r}$ is at least one.

For r = 2 the normalizing constant $c_r(u)$ is upper bounded by

$$c_r(u) = \sum_{v} d^{-r}(u, v) \le \sum_{k=1}^{2n} (4k)k^{-2} \le 4\sum_{k=1}^{2n} \frac{1}{k} = \theta(\ln n).$$

For r < 2, the normalizing constant $c_r(u)$ is lower bounded by

$$c_r(u) = \sum_{v} d^{-r}(u, v) \ge \sum_{k=1}^{n/2} (k) k^{-r} \ge \sum_{k=n/4}^{n/2} k^{1-r}.$$

The summation $\sum_{k=n/4}^{n/2} k^{1-r}$ has $\frac{n}{4}$ terms, the smallest of which is $\left(\frac{n}{4}\right)^{1-r}$ or $\left(\frac{n}{2}\right)^{1-r}$ depending on whether r is greater or less than one. This gives the following lower bound on $c_r(u)$.

$$c_r(u) \ge \frac{n}{4}\omega(n^{1-r}) = \omega(n^{2-r}).$$

No short paths exist for the r > 2 case.

For r > 2, we first show that for at least half of the pairs of vertices, there is no short path between them. We begin by showing that the expected number of edges of length greater than $n^{\frac{r+2}{2r}}$ goes to zero. The probability of an edge from u to v is $d^{-r}(u,v)/c_r(u)$ where $c_r(u)$ is lower bounded by a constant. Thus the probability that a particular edge of length greater than or equal to $n^{\frac{r+2}{2r}}$ is chosen is upper bounded by some contant ctimes $\left(n^{\frac{r+2}{2r}}\right)^{-r}$ or $cn^{-\left(\frac{r+2}{2}\right)}$. Since there are n^2 long edges, the expected number of edges of length at least $n^{\frac{r+2}{2r}}$ is at most $cn^2n^{-\frac{(r+2)}{2}}$ or $cn^{\frac{2-r}{2}}$, which for r > 2 goes to zero. Thus, by the first moment method, almost surely, there are no such edges. For at least half of the pairs of vertices, the grid distance, measured by grid edges between the vertices, is greater than or equal to n/4. Any path between them must have at least $\frac{1}{4}n/n^{\frac{r+2}{2r}} = \frac{1}{4}n^{\frac{r-2}{2r}}$ edges since there are no edges longer than $n^{\frac{r+2}{2r}}$ and so there is no polylog length path.

An algorithm for the r = 2 case

For r = 2, the local algorithm that selects the edge that ends closest to the destination t finds a path of expected length $O(\ln n)^3$. Suppose the algorithm is at a vertex u which is a at distance k from t. Then within an expected $O(\ln n)^2$ steps, the algorithm reaches a point at distance at most k/2. The reason is that there are $\Omega(k^2)$ vertices at distance at most k/2 from t. Each of these vertices is at distance at most k+k/2 = O(k) from u. See Figure 4.19. Recall that the normalizing constant c_r is upper bounded by $O(\ln n)$, and hence, the constant of proportionality is lower bounded by some constant times $1/\ln n$. Thus, the probability that the long-distance edge from u goes to one of these vertices is at least

$$\Omega(k^2 k^{-r} / \ln n) = \Omega(1 / \ln n).$$

Consider $\Omega(\ln n)^2$ steps of the path from u. The long-distance edges from the points visited at these steps are chosen independently and each has probability $\Omega(1/\ln n)$ of reaching within k/2 of t. The probability that none of them does is

$$\left(1 - \Omega(1/\ln n)\right)^{c(\ln n)^2} = c_1 e^{-\ln n} = \frac{c_1}{n}$$

for a suitable choice of constants. Thus, the distance to t is halved every $O(\ln n)^2$ steps and the algorithm reaches t in an expected $O(\ln n)^3$ steps.

A local algorithm cannot find short paths for the r < 2 case

For r < 2 no local polylog time algorithm exists for finding a short path. To illustrate the proof, we first give the proof for the special case r = 0, and then give the proof for r < 2.

When r = 0, all vertices are equally likely to be the end point of a long distance edge. Thus, the probability of a long distance edge hitting one of the *n* vertices that are within distance \sqrt{n} of the destination is 1/n. Along a path of length \sqrt{n} , the probability that the path does not encounter such an edge is $(1 - 1/n)^{\sqrt{n}}$. Now,

$$\lim_{n \to \infty} \left(1 - \frac{1}{n} \right)^{\sqrt{n}} = \lim_{n \to \infty} \left(1 - \frac{1}{n} \right)^{n \frac{1}{\sqrt{n}}} = \lim_{n \to \infty} e^{-\frac{1}{\sqrt{n}}} = 1.$$

Since with probability 1/2 the starting point is at distance at least n/4 from the destination and in \sqrt{n} steps, the path will not encounter a long distance edge ending within

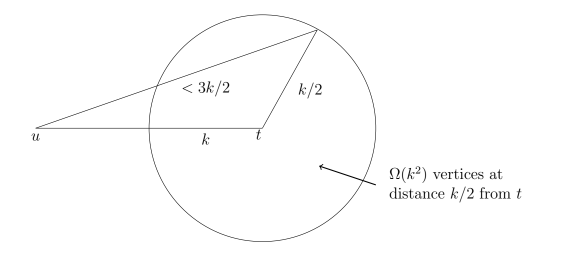


Figure 4.19: Small worlds.

distance \sqrt{n} of the destination, for at least half of the starting points the path length will be at least \sqrt{n} . Thus, the expected time is at least $\frac{1}{2}\sqrt{n}$ and hence not in polylog time.

For the general r < 2 case, we show that a local algorithm cannot find paths of length $O(n^{(2-r)/4})$. Let $\delta = (2-r)/4$ and suppose the algorithm finds a path with at most n^{δ} edges. There must be a long-distance edge on the path which terminates within distance n^{δ} of t; otherwise, the path would end in n^{δ} grid edges and would be too long. There are $O(n^{2\delta})$ vertices within distance n^{δ} of t and the probability that the long distance edge from one vertex of the path ends at one of these vertices is at most $n^{2\delta} \left(\frac{1}{n^{2-r}}\right) = n^{(r-2)/2}$. To see this, recall that the lower bound on the normalizing constant is $\theta(n^{2-r})$ and hence an upper bound on the probability that the long distance edge from one of the n^{δ} vertices on the path hits any one of the $n^{2\delta}$ vertices within distance n^{δ} of t is $n^{2\delta} \frac{1}{n^{2-r}} = n^{\frac{r-2}{2}}$. The probability that this happens for any one of the n^{δ} vertices on the path is at most $n^{\frac{r-2}{2}}n^{\delta} = n^{\frac{r-2}{2}/4} = n^{(r-2)/4} = o(1)$ as claimed.

Short paths exist for r < 2

Finally we show for r < 2 that there are $O(\ln n)$ length paths between s and t. The proof is similar to the proof of Theorem 4.17 showing $O(\ln n)$ diameter for G(n, p) when p is $\Omega(\ln n/n)$, so we do not give all the details here. We give the proof only for the case when r = 0.

For a particular vertex v, let S_i denote the set of vertices at distance i from v. Using only local edges, if i is $O(\sqrt{\ln n})$, then $|S_i|$ is $\Omega(\ln n)$. For later i, we argue a constant factor growth in the size of S_i as in Theorem 4.17. As long as $|S_1| + |S_2| + \cdots + |S_i| \le n^2/2$, for each of the $n^2/2$ or more vertices outside, the probability that the vertex is not in S_{i+1} is $(1 - \frac{1}{n^2})^{|S_i|} \leq 1 - \frac{|S_i|}{2n^2}$ since the long-distance edge from each vertex of S_i chooses a long-distance neighbor at random. So, the expected size of S_{i+1} is at least $|S_i|/4$ and using Chernoff, we get constant factor growth up to $n^2/2$. Thus, for any two vertices vand w, the number of vertices at distance $O(\ln n)$ from each is at least $n^2/2$. Any two sets of cardinality at least $n^2/2$ must intersect giving us a $O(\ln n)$ length path from v to w.

4.11 Bibliographic Notes

The G(n, p) random graph model is from Erdös Rényi [ER60]. Among the books written on properties of random graphs a reader may wish to consult Frieze and Karonski [FK15], Jansen, Luczak and Ruciński [JLR00],or Bollobás [Bol01]. Material on phase transitions can be found in [BT87]. The work on phase transitions for CNF was started by Chao and Franco [CF86]. Further work was done in [FS96], [AP03], [Fri99], and others. The proof here that the SC algorithm produces a solution when the number of clauses is cn for $c < \frac{2}{3}$ is from [Chv92].

For material on the giant component consult [Kar90] or [JKLP93]. Material on branching process can be found in [AN72]. The phase transition for giant components in random graphs with given degree distributions is from Molloy and Reed [MR95a].

There are numerous papers on growth models. The material in this chapter was based primarily on [CHK⁺] and [BA]. The material on small world is based on Kleinberg, [Kle00] which follows earlier work by Watts and Strogatz [WS98].

4.12 Exercises

Exercise 4.1 Search the World Wide Web to find some real world graphs in machine readable form or data bases that could automatically be converted to graphs.

- 1. Plot the degree distribution of each graph.
- 2. Compute the average degree of each graph.
- 3. Count the number of connected components of each size in each graph.
- 4. Describe what you find.
- 5. What is the average vertex degree in each graph? If the graph were a G(n, p) graph, what would the value of p be?
- 6. Spot differences between your graphs and G(n, p) for p from the last part. [Look at sizes of connected components, cycles, size of giant component.]

Exercise 4.2 In G(n,p) the probability of a vertex having degree k is $\binom{n}{k}p^k(1-p)^{n-k}$.

- 1. Show by direct calculation that the expected degree is np.
- 2. Compute directly the variance of the distribution.
- 3. Where is the mode of the binomial distribution for a given value of p? The mode is the point at which the probability is maximum.

Exercise 4.3

- 1. Plot the degree distribution for G(1000, 0.003).
- 2. Plot the degree distribution for G(1000, 0.030).

Exercise 4.4 To better understand the binomial distribution plot $\binom{n}{k}p^k(1-p)^{n-k}$ as a function of k for n = 50 and k = 0.05, 0.5, 0.95. For each value of p check the sum over all k to ensure that the sum is one.

Exercise 4.5 In $G(n, \frac{1}{n})$, argue that with high probability there is no vertex of degree greater than $\frac{6 \log n}{\log \log n}$ (i.e. ,the probability that such a vertex exists goes to zero as n goes to infinity). You may use the Poisson approximation and may wish to use the fact that $k! \geq (\frac{k}{\epsilon})^k$.

Exercise 4.6 The example of Section 4.1.1 showed that if the degrees in $G(n, \frac{1}{n})$ were independent there would almost surely be a vertex of degree $\Omega(\log n / \log \log n)$. However, the degrees are not independent. Show how to overcome this difficulty.

Exercise 4.7 Let f(n) be a function that is asymptotically less than n. Some such functions are 1/n, a constant d, $\log n$ or $n^{\frac{1}{3}}$. Show that

$$\left(1 + \frac{f(n)}{n}\right)^n \simeq e^{f(n)(1 \pm o(1))}.$$

for large n. That is

$$\lim_{n \to \infty} \frac{\ln\left[\left(1 + \frac{f(n)}{n}\right)^n\right]}{f(n)} = 1.$$

Exercise 4.8

- 1. In the limit as n goes to infinity, how does $\left(1-\frac{1}{n}\right)^{n\ln n}$ behave.
- 2. What is $\lim_{n \to \infty} \left(\frac{n+1}{n}\right)^n$?

Exercise 4.9 Consider a random permutation of the integers 1 to n. The integer i is said to be a fixed point of the permutation if i is the integer in the i^{th} position of the permutation. Use indicator variables to determine the expected number of fixed points in a random permutation.

Exercise 4.10 Generate a graph $G(n, \frac{d}{n})$ with n = 1000 and d=2, 3, and 6. Count the number of triangles in each graph. Try the experiment with n=100.

Exercise 4.11 What is the expected number of squares (4-cycles) in $G\left(n, \frac{d}{n}\right)$? What is the expected number of 4-cliques in $G\left(n, \frac{d}{n}\right)$?

Exercise 4.12 Carry out an argument, similar to the one used for triangles, to show that $p = \frac{1}{n^{2/3}}$ is a threshold for the existence of a 4-clique. A 4-clique consists of four vertices with all $\binom{4}{2}$ edges present.

Exercise 4.13 What is the expected number of simple paths of length 3, $\log n$, \sqrt{n} , and n-1 in $G(n, \frac{d}{n})$? A simple path is a path where no vertex appears twice as in a cycle. The expected number of simple paths of a given length being infinite does not imply that a graph selected at random has such a path.

Exercise 4.14 Let x be an integer chosen uniformly at random from $\{1, 2, ..., n\}$. Count the number of distinct prime factors of n. The exercise is to show that the number of prime factors almost surely is $\Theta(\ln \ln n)$. Let p stand for a prime number between 2 and n.

1. For each fixed prime p, let I_p be the indicator function of the event that p divides x. Show that $E(I_p) = \frac{1}{p} + O\left(\frac{1}{n}\right)$.

- 2. The random variable of interest, $y = \sum_{p} I_p$, is the number of prime divisors of x picked at random. Show that the variance of y is $O(\ln \ln n)$. For this, assume the known result that the number of primes p between 2 and n is $O(n/\ln n)$ and that $\sum_{p} \frac{1}{p} = \ln \ln n$. To bound the variance of y, think of what $E(I_pI_q)$ is for $p \neq q$, both primes.
- 3. Use (1) and (2) to prove that the number of prime factors is almost surely $\theta(\ln \ln n)$.

Exercise 4.15 Suppose one hides a clique of size k in a random graph $G(n, \frac{1}{2})$. I.e., in the random graph, choose some subset S of k vertices and put in the missing edges to make S a clique. Presented with the modified graph, find S. The larger S is, the easier it should be to find. In fact, if k is more than $c\sqrt{n \ln n}$, then with high probability the clique leaves a telltale sign identifying S as the k vertices of largest degree. Prove this statement by appealing to Theorem 4.1. It remains a puzzling open problem to do this when k is smaller, say, $O(n^{1/3})$.

Exercise 4.16 The clique problem in a graph is to find the maximal size clique. This problem is known to be NP-hard and so a polynomial time algorithm is thought unlikely. We can ask the corresponding question about random graphs. For example, in $G(n, \frac{1}{2})$ there almost surely is a clique of size $(2 - \varepsilon) \log n$ for any $\varepsilon > 0$. But it is not known how to find one in polynomial time.

- 1. Show that in $G(n, \frac{1}{2})$, there almost surely are no cliques of size $\geq 2\log_2 n$.
- 2. Use the second moment method to show that in $G(n, \frac{1}{2})$, almost surely there are cliques of size $(2 \varepsilon) \log_2 n$.
- 3. Show that for any $\varepsilon > 0$, a clique of size $(2 \varepsilon) \log n$ can be found in $G\left(n, \frac{1}{2}\right)$ in time $n^{O(\ln n)}$ if one exists.
- 4. Give an $O(n^2)$ algorithm that finds a clique of size $\Omega(\log n)$ in $G(n, \frac{1}{2})$ with high probability. Hint: use a greedy algorithm. Apply your algorithm to $G(1000, \frac{1}{2})$. What size clique do you find?
- 5. An independent set in a graph is a set of vertices such that no two of them are connected by an edge. Give a polynomial time algorithm for finding an independent set in $G(n, \frac{1}{2})$ of size $\Omega(\log n)$ with high probability.

Exercise 4.17 Suppose H is a fixed graph on cn vertices with $\frac{1}{4}c^2(\log n)^2$ edges. Show that if $c \ge 2$, with high probability, H does not occur as a subgraph of G(n, 1/4).

Exercise 4.18 Given two instances, G_1 and G_2 of $G(n, \frac{1}{2})$, what is the largest vertexinduced subgraph common to both G_1 and G_2 ? **Exercise 4.19** (Birthday problem) What is the number of integers that must be drawn with replacement from a set of n integers so that some integer, almost surely, will be selected twice?

Exercise 4.20 Suppose you have an algorithm for finding communities in a social network. Assume that the way the algorithm works is that given the graph G for the social network, it finds a subset of vertices satisfying a desired property P (the specifics of property P are unimportant for this question). If there are multiple subsets S of vertices that satisfy property P, assume that the algorithm finds one such set S at random.

In running the algorithm you find thousands of communities and wonder how many communities there are in the graph. Finally, when you find the $10,000^{th}$ community, it is a duplicate. It is the same community as one found earlier. Use the birthday problem to derive an estimate of the total number of communities in G.

Exercise 4.21 Do a breadth first search in $G(n, \frac{d}{n})$ with d > 1 starting from some vertex. The number of discovered vertices, z_i , after i steps has distribution $Binomial(n, p_i)$ where $p_i = 1 - (1 - \frac{d}{n})^i$. If the connected component containing the start vertex has i vertices, then $z_i = i$. Show that as $n \to \infty$ (and d is a fixed constant), $Prob(z_i = i) \in o(1/n)$ unless $i \leq c_1 \ln n$ or $i \geq c_2 n$ for some constants c_1, c_2 .

Exercise 4.22 For $f(x) = 1 - e^{-dx} - x$, what is the value of $x_{max} = \arg \max f(x)$? What is the value of $f(x_{max})$? Recall from the text that in a breadth first search of $G(n, \frac{d}{n})$, f(x) is the expected normalized size of the frontier (size of frontier divided by n) at normalized time x (x = t/n). Where does the maximum expected value of the frontier of a breadth search in $G(n, \frac{d}{n})$ occur as a function of n?

Exercise 4.23 If y and z are independent, nonnegative, integer valued random variables, then the generating function of the sum y + z is the product of the generating function of y and z. Show that this follows from $E(x^{y+z}) = E(x^yx^z) = E(x^y)E(x^z)$.

Exercise 4.24 Let $f_j(x)$ be the j^{th} iterate of the generating function f(x) of a branching process. When m > 1, $\lim_{j\to\infty} f_j(x) = q$ for 0 < x < 1. In the limit this implies $Prob(z_j = 0) = q$ and $Prob(z_j = i) = 0$ for all nonzero finite values of i. Shouldn't the probabilities add up to 1? Why is this not a contradiction?

Exercise 4.25 Try to create a probability distribution for a branching process which varies with the current population in which future generations neither die out, nor grow to infinity.

Exercise 4.26 Let d be a constant strictly greater than 1. Show that for a branching process with number of children distributed as $Binomial(n - c_1 n^{2/3}, \frac{d}{n})$, the root of the f(x) = 1 in (0, 1) is at most a constant strictly less than 1.

Exercise 4.27 Randomly generate G(50, p) for several values of p. Start with $p = \frac{1}{50}$.

- 1. For what value of p do cycles first appear?
- 2. For what value of p do isolated vertices disappear and the graphs become connected?

Exercise 4.28 Consider G(n,p) with $p = \frac{1}{3n}$.

- 1. Use the second moment method to show that with high probability there exists a simple path of length 10.
- 2. Argue that on the other hand, it is unlikely there exists any cycle of length 10.

Exercise 4.29 Complete the second moment argument of Theorem 4.13 to show that for $p = \frac{d}{n}$, d > 1, G(n, p) almost surely has a cycle.

Hint: If two cycles share one or more edges, then the union of the two cycles is at least one greater than the union of the vertices.

Exercise 4.30 Draw a tree with 10 vertices and label each vertex with a unique integer from 1 to 10. Construct the Prüfer sequence (Appendix 12.9.6) for the tree. Given the Prüfer sequence, recreate the tree.

Exercise 4.31 Construct the tree corresponding to the following Prüfer sequences (Appendix 12.9.6)

- 1. 113663 (1,2), (1,3), (1,4), (3,5), (3,6), (6,7), and (6,8)
- 2. 552833226.

Exercise 4.32 What is the expected number of isolated vertices in G(n,p) for $p = \frac{1}{2} \frac{\ln n}{n}$?

Exercise 4.33 Theorem 4.17 shows that for some c > 0 and $p = c \ln n/n$, G(n, p) has diameter $O(\ln n)$. Tighten the argument to pin down as low a value as possible for c.

Exercise 4.34 What is diameter of G(n,p) for various values of p?

Exercise 4.35

- 1. List five increasing properties of G(n, p).
- 2. List five non increasing properties.

Exercise 4.36 Consider generating the edges of a random graph by flipping two coins, one with probability p_1 of heads and the other with probability p_2 of heads. Add the edge to the graph if either coin comes down heads. What is the value of p for the generated G(n, p) graph?

Exercise 4.37 In the proof of Theorem 4.19, we proved for $p_0(n)$ such that $\lim_{n\to\infty} \frac{p_0(n)}{p(n)} = 0$ that $G(n, p_0)$ almost surely did not have property Q. Give the symmetric argument that for any $p_1(n)$ such that $\lim_{n\to\infty} \frac{p(n)}{p_1(n)} = 0$, $G(n, p_1)$ almost surely has property Q.

Exercise 4.38 Consider a model of a random subset N(n, p) of integers $\{1, 2, ..., n\}$ defined by independently at random including each of $\{1, 2, ..., n\}$ into the set with probability p. Define what an "increasing property" of N(n, p) means. Prove that every increasing property of N(n, p) has a threshold.

Exercise 4.39 N(n,p) is a model of a random subset of integers $\{1, 2, ..., n\}$ defined by independently at random including each of $\{1, 2, ..., n\}$ into the set with probability p. What is the threshold for N(n,p) to contain

- 1. a perfect square,
- 2. a perfect cube,
- 3. an even number,
- 4. three numbers such that x + y = z?

Exercise 4.40 Explain why the property that N(n, p) contains the integer 1 has a threshold. What is the threshold?

Exercise 4.41 The Sudoku game consists of a 9×9 array of squares. The array is partitioned into nine 3×3 squares. Each small square should be filled with an integer between 1 and 9 so that each row, each column, and each 3×3 square contains exactly one copy of each integer. Initially the board has some of the small squares filled in in such a way that there is exactly one way to complete the assignments of integers to squares. Some simple rules can be developed to fill in the remaining squares such as if a row does not contain a given integer and if every column except one in which the square in the row is blank contains the integer, then place the integer in the remaining blank square in the row. Explore phase transitions for the Sudoku game. Some possibilities are:

- 1. Start with a 9 × 9 array of squares with each square containing a number between 1 and 9 such that no row, column, or 3 × 3 square has two copies of any integer. Develop a set of simple rules for filling in squares such as if a row does not contain a given integer and if every column except one in which the square in the row is blank contains the integer, then place the integer in the remaining blank entry in the row. How many integers can you randomly erase and your rules will still completely fill in the board?
- 2. Generalize the Sudoku game for arrays of size $n^2 \times n^2$. Develop a simple set of rules for completing the game. Start with a legitimate completed array and erase k entries at random. Experimentally determine the threshold for the integer k such that if only k entries of the array are erased, your set of rules will find a solution?

Exercise 4.42 In a square $n \times n$ grid, each of the $O(n^2)$ edges is randomly chosen to be present with probability p and absent with probability 1 - p. Consider the increasing property that there is a path from the bottom left corner to the top right corner which always goes to the right or up. Show that p = 1/2 is a threshold for the property. Is it a sharp threshold?

Exercise 4.43 The threshold property seems to be related to uniform distributions. What if we considered other distributions? Consider a model where i is selected from the set $\{1, 2, ..., n\}$ with probability $\frac{c(n)}{i}$. Is there a threshold for perfect squares? Is there a threshold for arithmetic progressions?

Exercise 4.44 Modify the proof that every increasing property of G(n, p) has a threshold to apply to the 3-CNF satisfiability problem.

Exercise 4.45 Evaluate $\left(1-\frac{1}{2^k}\right)^{2^k}$ for k=3, 5, and 7. How close is it to 1/e?

Exercise 4.46 Randomly generate clauses for a Boolean formula in 3-CNF. Compute the number of solutions and the number of connected components of the solution set as a function of the number of clauses generated. What happens?

Exercise 4.47 Consider a random process for generating a Boolean function f in conjunctive normal form where each of c clauses is generated by placing each of n variables in the clause with probability p and complementing the variable with probability $\frac{1}{2}$. What is the distribution of clause sizes for various p such as $p = \frac{3}{n}$, $\frac{1}{2}$, other values? Experimentally determine the threshold value of p for f to cease to be satisfied.

Exercise 4.48 For a random 3-CNF formula with n variables and cn clauses, what is the expected number of satisfying assignments?

Exercise 4.49 Which of the following variants of the SC algorithm admit a theorem like Theorem 4.21?

- 1. Among all clauses of least length, pick the first one in the order in which they appear in the formula.
- 2. Set the literal appearing in most clauses independent of length to 1.

Exercise 4.50 Suppose we have a queue of jobs serviced by one server. There is a total of n jobs in the system. At time t, each remaining job independently decides to join the queue to be serviced with probability p = d/n, where d < 1 is a constant. Each job has a processing time of 1 and at each time the server services one job, if the queue is nonempty. Show that with high probability, no job waits more than $\Omega(\ln n)$ time to be serviced once it joins the queue.

Exercise 4.51 Consider G(n, p). Where is the phase transition for 2-colorability? Hint: For p = d/n with d < 1, G(n, p) is acyclic, so it is bipartite and hence 2-colorable. When $pn \to \infty$, the expected number of triangles goes to infinity. Show that, almost surely, there is a triangle. What does this do for 2-colorability?

Exercise 4.52 A vertex cover of size k for a graph is a set of k vertices such that one end of each edge is in the set. Experimentally play with the following problem. For $G(n, \frac{1}{2})$, for what value of k is there a vertex cover of size k?

Exercise 4.53 Consider graph 3-colorability. Randomly generate the edges of a graph and compute the number of solutions and the number of connected components of the solution set as a function of the number of edges generated. What happens?

Exercise 4.54 Construct an example of a formula which is satisfiable, but the SC heuristic fails to find a satisfying assignment.

Exercise 4.55 In G(n, p), let x_k be the number of connected components of size k. Using x_k , write down the probability that a randomly chosen vertex is in a connected component of size k. Also write down the expected size of the connected component containing a randomly chosen vertex.

Exercise 4.56 In a G(n,p) graph with p asymptotically greater than $\frac{1}{n}$, show that $\sum_{i=0}^{\infty} i(i-2)\lambda_i > 0$ where λ_i is the fraction of vertices of degree i.

Exercise 4.57 Describe several methods of generating a random graph with a given degree distribution. Describe differences in the graphs generated by the different methods.

Exercise 4.58 Consider generating a random graph adding one edge at a time. Let n(i,t) be the number of components of size i at time t.

$$n(1,1) = n$$

$$n(1,t) = 0 \quad t > 1$$

$$n(i,t) = n(i,t-1) + \sum \frac{j(i-j)}{n^2} n(j,t-1) n(i-j,t-1) - \frac{2i}{n} n(i)$$

Compute n(i,t) for a number of values of i and t. What is the behavior? What is the sum of n(i,t) for fixed t and all i? Can you write a generating function for n(i,t)?

Exercise 4.59 The global clustering coefficient of a graph is defined as follows. Let d_v be the degree of vertex v and let e_v be the number of edges connecting pairs of vertices that are adjacent to vertex v. The global clustering coefficient c is given by

$$c = \sum_{v} \frac{2e_v}{d_v(d_v-1)}.$$

In a social network, for example, it measures what fraction of pairs of friends of each person are themselves friends. If many are, the clustering coefficient is high. What is c for a random graph with $p = \frac{d}{n}$ in the limit as n goes to infinity? For a denser graph? Compare this value to that for some social network.

Exercise 4.60 Consider a structured graph, such as a grid or cycle, and gradually add edges or reroute edges at random. Let L be the average distance between all pairs of vertices in a graph and let C be the ratio of triangles to connected sets of three vertices. Plot L and C as a function of the randomness introduced.

Exercise 4.61 Consider an $n \times n$ grid in the plane.

- 1. Prove that for any vertex u, there are at least k vertices at distance k for $1 \le k \le n/2$.
- 2. Prove that for any vertex u, there are at most 4k vertices at distance k.
- 3. Prove that for one half of the pairs of points, the distance between them is at least n/4.

Exercise 4.62 Recall the definition of a small-world graph in Section 4.10. Show that in a small-world graph with $r \leq 2$, that there exist short paths with high probability. The proof for r = 0 is in the text.

Exercise 4.63 Change the small worlds graph as follows. Start with a $n \times n$ grid where each vertex has one long-distance edge to a vertex chosen uniformly at random. These are exactly like the long-distance edges for r = 0. But the grid edges are not present. Instead, we have some other graph with the property that for each vertex, there are $\Theta(t^2)$ vertices at distance t from the vertex for $t \leq n$. Show that, almost surely, the diameter is $O(\ln n)$.

Exercise 4.64 Consider an n-node directed graph with two random out-edges from each node. For two vertices s and t chosen at random, prove that with high probability there exists a path of length at most $O(\ln n)$ from s to t.

Exercise 4.65 Explore the concept of small world by experimentally determining the answers to the following questions:

- 1. How many edges are needed to disconnect a small world graph? By disconnect we mean at least two pieces each of reasonable size. Is this connected to the emergence of a giant component?
- 2. How does the diameter of a graph consisting of a cycle change as one adds a few random long distance edges?

Exercise 4.66 In the small world model, would it help if the algorithm could look at edges at any node at a cost of one for each node looked at?

Exercise 4.67 Consider the $n \times n$ grid in the section on small world graphs. If the probability of an edge from vertex u to vertex v is proportional to $d^{-r}(u, v)$, show that the constant of proportionality $c_r(u)$ is

$$\begin{array}{ll} \theta(n^{2-r}) & for \ r > 2 \\ \theta(\ln n) & for \ r = 2 \\ \theta(1) & for \ r < 2 \end{array}$$

Exercise 4.68 Show that for r < 2 in the small world graph model that short paths exist but a polylog length path is unlikely to encounter a long distance edge whose end point is close to the destination.

Exercise 4.69 Make a list of the ten most interesting things you learned about random graphs.

5 Random Walks and Markov Chains

A random walk on a directed graph consists of a sequence of vertices generated from a start vertex by selecting an edge, traversing the edge to a new vertex, and repeating the process. If the graph is strongly connected, then the fraction of time the walk spends at the vertices of the graph converges to a stationary probability distribution.

Since the graph is directed, there might be vertices with no out edges and hence nowhere for the walk to go. Vertices in a strongly connected component with no in edges from the remainder of the graph can never be reached unless the component contains the start vertex. Once a walk leaves a strongly connected component, it can never return. Most of our discussion of random walks will involve strongly connected graphs.

Start a random walk at a vertex x_0 and think of the starting probability distribution as putting a mass of one on x_0 and zero on every other vertex. More generally, one could start with any probability distribution \mathbf{p} , where \mathbf{p} is a row vector with nonnegative components summing to one, with p_x being the probability of starting at vertex x. The probability of being at vertex x at time t + 1 is the sum over each adjacent vertex y of being at y at time t and taking the transition from y to x. Let \mathbf{p}_t be a row vector with a component for each vertex specifying the probability mass of the vertex at time t and let \mathbf{p}_{t+1} be the row vector of probabilities at time t + 1. In matrix notation¹⁵

$\mathbf{p_t} P = \mathbf{p_{t+1}}$

where the ij^{th} entry of the matrix P is the probability of the walk at vertex i selecting the edge to vertex j.

A fundamental property of a random walk is that in the limit, the long-term average probability of being at a particular vertex is independent of the start vertex, or an initial probability distribution over vertices, provided only that the underlying graph is strongly connected. The limiting probabilities are called the *stationary probabilities*. This fundamental theorem is proved in the next section.

A special case of random walks, namely random walks on undirected graphs, has important connections to electrical networks. Here, each edge has a parameter called *conductance*, like electrical conductance. If the walk is at vertex v, it chooses an edge from among all edges incident to u to walk to the next vertex with probability proportional to its conductance. Certain basic quantities associated with random walks are hitting time, the expected time to reach vertex y starting at vertex x, and cover time, the expected time to visit every vertex. Qualitatively, these quantities are all bounded above by polynomials in the number of vertices. The proofs of these facts will rely on the analogy between random walks and electrical networks.

¹⁵Probability vectors are represented by row vectors to simplify notation in equations like the one here.

random walk	Markov chain
graph	stochastic process
vertex	state
strongly connected	persistent
aperiodic	aperiodic
strongly connected and aperiodic	ergotic
undirected graph	time reversible

Table 5.1: Correspondence between terminology of random walks and Markov chains

Aspects of the theory of random walks were developed in computer science with an important application in defining the pagerank of pages on the World Wide Web by their stationary probability. An equivalent concept called a *Markov chain* had previously been developed in the statistical literature. A Markov chain has a finite set of *states*. For each pair of states x and y, there is a *transition probability* p_{xy} of going from state x to state y where for each x, $\sum_{y} p_{xy} = 1$. A random walk in the Markov chain starts at some state. At a given time step, if it is in state x, the next state y is selected randomly with probability p_{xy} . A Markov chain can be represented by a directed graph with a vertex representing each state and an edge with weight p_{xy} from vertex x to vertex y. We say that the Markov chain is *connected* if the underlying directed graph is strongly connected. That is, if there is a directed path from every vertex to every other vertex. The matrix P consisting of the p_{xy} is called the *transition probability matrix* of the chain. The terms "random walk" and "Markov chain" are used interchangeably. The correspondence between the terminologies of random walks and Markov chains is given in Table 5.1.

A state of a Markov chain is *persistent* if it has the property that should the state ever be reached, the random process will return to it with probability one. This is equivalent to the property that the state is in a strongly connected component with no out edges. For most of the chapter, we assume that the underlying directed graph is strongly connected. We discuss here briefly what might happen if we do not have strong connectivity. Consider the directed graph in Figure 5.1b with three strongly connected components, A, B, and C. Starting from any vertex in A, there is a nonzero probability of eventually reaching any vertex in A. However, the probability of returning to a vertex in A is less than one and thus vertices in A, and similarly vertices in B, are not persistent. From any vertex in C, the walk eventually will return with probability one to the vertex, since there is no way of leaving component C. Thus, vertices in C are persistent.

A connected Markov Chain is said to be *aperiodic* if the greatest common divisor of the lengths of directed cycles is one. It is known that for connected aperiodic chains, the probability distribution of the random walk converges to a unique stationary distribu-

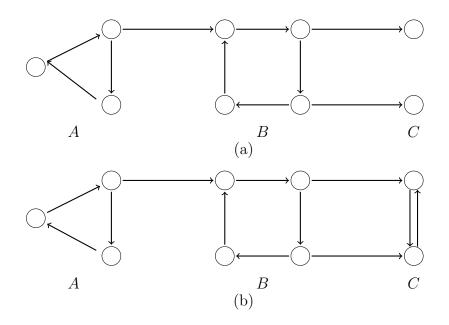


Figure 5.1: (a) A directed graph with vertices having no out out edges and a strongly connected component A with no in edges.

(b) A directed graph with three strongly connected components.

tion. Aperiodicity is a technical condition needed in this proof. Here, we do not prove this theorem and do not worry about aperiodicity at all. It turns out that if we take the average probability distribution of the random walk over the first t steps, then this average converges to a limiting distribution for connected chains (without assuming aperiodicity) and this average is what one uses in practice. We do prove this limit theorem and explain its uses in what is called the "Markov Chain Monte Carlo (MCMC)" method.

Markov chains are used to model situations where all the information of the system necessary to predict the future can be encoded in the current state. A typical example is speech, where for a small k the current state encodes the last k syllables uttered by the speaker. Given the current state, there is a certain probability of each syllable being uttered next and these can be used to calculate the transition probabilities. Another example is a gambler's assets, which can be modeled as a Markov chain where the current state is the amount of money the gambler has on hand. The model would only be valid if the gambler's bets depend only on current assets, not the past history.

Later in the chapter, we study the widely used Markov Chain Monte Carlo method (MCMC). Here, the objective is to sample a large space according to some probability distribution p. The number of elements in the space may be very large, say 10^{100} . One designs a Markov chain where states correspond to the elements of the space. The transition probabilities of the chain are designed so that the stationary probability of the chain is the probability distribution p with which we want to sample. One samples by taking

a random walk until the probability distribution is close to the stationary distribution of the chain and then selects the point the walk is at. The walk continues a number of steps until the probability distribution is no longer dependent on where the walk was when the first element was selected. A second point is then selected, and so on. Although it is impossible to store the graph in a computer since it has 10^{100} vertices, to do the walk one needs only store the vertex the walk is at and be able to generate the adjacent vertices by some algorithm. What is critical is that the probability of the walk converges to the stationary probability in time logarithmic in the number of states.

We mention two motivating examples. The first is to select a point at random in d-space according to a probability density such as a Gaussian. Put down a grid and let each grid point be a state of the Markov chain. Given a probability density p, design transition probabilities of a Markov chain so that the stationary distribution is p. In general, the number of states grows exponentially in the dimension d, but if the time to converge to the stationary distribution grows polynomially in d, then one can do a random walk on the graph until convergence to the stationary probability. Once the stationary probability has been reached, one selects a point. To select a set of points, one must walk a number of steps between each selection so that the probability of the current point is independent of the previous point. By selecting a number of points one can estimate the probability of a region by observing the number of selected points in the region.

A second example is from physics. Consider an $n \times n$ grid in the plane with a particle at each grid point. Each particle has a spin of ± 1 . A configuration is a n^2 dimensional vector $\mathbf{v} = (v_1, v_2, \ldots, v_{n^2})$, where, v_t is the spin of the t^{th} particle. There are 2^{n^2} spin configurations. The energy of a configuration is a function $f(\mathbf{v})$ of the configuration, not of any single spin. A central problem in statistical mechanics is to sample spin configurations according to their probability. It is easy to design a Markov chain with one state per spin configuration so that the stationary probability of a state is proportional to the state's energy. If a random walk gets close to the stationary probability in time polynomial in n rather than 2^{n^2} , then one can sample spin configurations according to their probability.

The Markov Chain has 2^{n^2} states, one per configuration. Two states in the Markov chain are adjacent if and only if the corresponding configurations \mathbf{v} and \mathbf{u} differ in just one coordinate ($u_t = v_t$ for all but one t). The Metropilis-Hastings random walk, described in more detail in Section 5.2, has a transition probability from a configuration \mathbf{v} to an adjacent configuration \mathbf{u} of

$$\frac{1}{n^2}$$
 Min $\left(1, \frac{f(\mathbf{u})}{f(\mathbf{v})}\right)$.

As we will see, the Markov Chain has a stationary probability proportional to the energy. There are two more crucial facts about this chain. The first is that to execute a step in the chain, we do not need any global information of the whole chain, just the ratio $\frac{f(u)}{f(v)}$ is needed. The second is that under suitable assumptions, the chain approaches stationarity

in time polynomial in n.

A quantity called the *mixing time*, loosely defined as the time needed to get close to the stationary distribution, is often much smaller than the number of states. In Section 5.4, we relate the mixing time to a combinatorial notion called *normalized conductance* and derive upper bounds on the mixing time in several cases.

5.1 Stationary Distribution

Let \mathbf{p}_t be the probability distribution after t steps of a random walk. Define the long-term average probability distribution \mathbf{a}_t by

$$\mathbf{a}_{\mathbf{t}} = \frac{1}{t} \left(\mathbf{p}_{\mathbf{0}} + \mathbf{p}_{\mathbf{1}} + \dots + \mathbf{p}_{\mathbf{t}-\mathbf{1}} \right).$$

The fundamental theorem of Markov chains asserts that for a connected Markov chain, \mathbf{a}_t converges to a limit probability vector \mathbf{x} , which satisfies the equations $\mathbf{x}P = \mathbf{x}$; $\sum_i x_i = 1$ which we can rewrite as

$$\mathbf{x}[P-I,\mathbf{1}] = [\mathbf{0},1].$$

We will prove now that the matrix $[P-I, \mathbf{1}]$ has rank *n* provided the Markov Chain is connected. This implies that there is a unique solution to the equations $\mathbf{x}[P-I, \mathbf{1}] = [\mathbf{0}, \mathbf{1}]$. We denote this solution by $\boldsymbol{\pi}$. It has non-negative components and so is a probability vector. Since $\boldsymbol{\pi}P = \boldsymbol{\pi}$, we have that running one step of the Markov Chain starting with distribution $\boldsymbol{\pi}$ leaves us in the same distribution. Thus also running any number of steps of the Markov Chain starting the first step with $\boldsymbol{\pi}$ leaves the distribution still the same. For this reason, $\boldsymbol{\pi}$ is called the *stationary distribution*.

Lemma 5.1 Let P be the transition probability matrix for a connected Markov chain. The $n \times (n+1)$ matrix A = [P - I, 1] obtained by augmenting the matrix P - I with an additional column of ones has rank n.

Proof: If the rank of $A = [P - I, \mathbf{1}]$ was less than n there would be two linearly independent solutions to $A\mathbf{x} = \mathbf{0}$. Each row in P sums to one so each row in P - I sums to zero. Thus $(\mathbf{1}, 0)$, where all but the last coordinate is 1, is one solution. Assume there was a second solution (\mathbf{x}, α) perpendicular to $(\mathbf{1}, 0)$. Then $(P - I)\mathbf{x} + \alpha \mathbf{1} = \mathbf{0}$. Thus for each i, $\sum_{j} p_{ij}x_j - x_i + \alpha = 0$ or $x_i = \sum_{j} p_{ij}x_j + \alpha$. Each x_i is a convex combination of some of the x_j plus α . Since x is perpendicular to $\mathbf{1}$, not all x_i can be equal. Let $S = \{i : x_i = \operatorname{Max}_{j=1}^n x_j\}$ be the set of i for which x_i attains its maximum value. \overline{S} is not empty. Connectedness implies that there is some edge $(k, l), k \in S, l \in \overline{S}$. Thus, $x_k > \sum_j p_{kj} x_j$. Therefore α must be greater than 0 in $x_k = \sum_j p_{kj} x_j + \alpha$. A symmetric argument with the set of i with x_i taking its minimum value implies $\alpha < 0$ producing a contradiction proving the lemma.

Theorem 5.2 (Fundamental Theorem of Markov Chains) Let P be the transition probability matrix for a connected Markov chain, $\mathbf{p_t}$ the probability distribution after t steps of a random walk, and

$$\mathbf{a_t} = \frac{1}{t} \left(\mathbf{p_0} + \mathbf{p_1} + \dots + \mathbf{p_{t-1}} \right)$$

the long term average probability distribution. Then there is a unique probability vector $\boldsymbol{\pi}$ satisfying $\boldsymbol{\pi} P = \boldsymbol{\pi}$. Moreover, for any starting distribution, $\lim_{t\to\infty} \mathbf{a}_t$ exists and equals $\boldsymbol{\pi}$.

Proof: Note that \mathbf{a}_t is itself a probability vector, since its components are nonnegative and sum to one. After one step, the distribution of the Markov chain starting from the distribution \mathbf{a}_t is $\mathbf{a}_t P$. The change in probabilities due to this step is

$$\mathbf{a}_{\mathbf{t}}P - \mathbf{a}_{\mathbf{t}} = \frac{1}{t} \left[\mathbf{p}_{\mathbf{0}}P + \mathbf{p}_{\mathbf{1}}P + \dots + \mathbf{p}_{\mathbf{t}-\mathbf{1}}P \right] - \frac{1}{t} \left[\mathbf{p}_{\mathbf{0}} + \mathbf{p}_{\mathbf{1}} + \dots + \mathbf{p}_{\mathbf{t}-\mathbf{1}} \right]$$
$$= \frac{1}{t} \left[\mathbf{p}_{\mathbf{1}} + \mathbf{p}_{\mathbf{2}} + \dots + \mathbf{p}_{\mathbf{t}} \right] - \frac{1}{t} \left[\mathbf{p}_{\mathbf{0}} + \mathbf{p}_{\mathbf{1}} + \dots + \mathbf{p}_{\mathbf{t}-\mathbf{1}} \right]$$
$$= \frac{1}{t} \left(\mathbf{p}_{\mathbf{t}} - \mathbf{p}_{\mathbf{0}} \right).$$

Thus, $\mathbf{b_t} = \mathbf{a_t}P - \mathbf{a_t}$ satisfies $|\mathbf{b_t}| \le \frac{2}{t}$ and goes to zero as $t \to \infty$.

By Lemma (5.1) $A = [P - I, \mathbf{1}]$ has rank n. Since the first n columns of A sum to zero, the $n \times n$ submatrix B of A consisting of all its columns except the first is invertible. Let \mathbf{c}_t be obtained from $\mathbf{b}_t = \mathbf{a}_t P - \mathbf{a}_t$ by removing the first entry so that $\mathbf{a}_t B = [\mathbf{c}_t, 1]$. Then $\mathbf{a}_t = [\mathbf{c}_t, 1]B^{-1}$. Since $\mathbf{b}_t \to \mathbf{0}$, $[\mathbf{c}_t, 1] \to [\mathbf{0}, 1]$ and $\mathbf{a}_t \to [\mathbf{0}, 1]B^{-1}$. Denote $[\mathbf{0}, 1]B^{-1}$ as π . Since $\mathbf{a}_t \to \pi$, we have that π is a probability vector. Since $\mathbf{a}_t[P - I] = \mathbf{b}_t \to \mathbf{0}$, we get $\pi[P - I] = \mathbf{0}$. Since A has rank n, this is the unique solution as required.

Observe that the expected time r_x for a Markov chain starting in state x to return to state x is the reciprocal of the stationary probability of x. That is $r_x = \frac{1}{\pi_x}$. Intuitively this follows by observing that if a long walk always returns to state x in exactly r_x steps, the frequency of being in a state x would be $\frac{1}{r_x}$. A rigorous proof requires the Strong Law of Large Numbers.

We finish this section with the following lemma useful in establishing that a probability distribution is the stationary probability distribution for a random walk on a connected graph with edge probabilities.

Lemma 5.3 For a random walk on a strongly connected graph with probabilities on the edges, if the vector $\boldsymbol{\pi}$ satisfies $\pi_x p_{xy} = \pi_y p_{yx}$ for all x and y and $\sum_x \pi_x = 1$, then $\boldsymbol{\pi}$ is the stationary distribution of the walk.

Proof: Since π satisfies $\pi_x p_{xy} = \pi_y p_{yx}$, summing both sides, $\pi_x = \sum_y \pi_y p_{yx}$ and hence π satisfies $\pi = \pi P$. By Theorem 5.2, π is the unique stationary probability.

5.2 Markov Chain Monte Carlo

The Markov Chain Monte Carlo (MCMC) method is a technique for sampling a multivariate probability distribution $p(\mathbf{x})$, where $\mathbf{x} = (x_1, x_2, \ldots, x_d)$. The MCMC method is used to estimate the expected value of a function $f(\mathbf{x})$

$$E(f) = \sum_{\mathbf{x}} f(\mathbf{x}) p(\mathbf{x}).$$

If each x_i can take on two or more values, then there are at least 2^d values for \mathbf{x} , so an explicit summation requires exponential time. Instead, one could draw a set of samples, each sample \mathbf{x} with probability $p(\mathbf{x})$. Averaging f over these samples provides an estimate of the sum.

To sample according to $p(\mathbf{x})$, design a Markov Chain whose states correspond to the possible values of \mathbf{x} and whose stationary probability distribution is $p(\mathbf{x})$. There are two general techniques to design such a Markov Chain: the Metropolis-Hastings algorithm and Gibbs sampling, which we will describe in the next section. The Fundamental Theorem of Markov Chains, Theorem 5.2, states that the average of f over states seen in a sufficiently long run is a good estimate of E(f). The harder task is to show that the number of steps needed before the long-run average probabilities are close to the stationary distribution grows polynomially in d, though the total number of states may grow exponentially in d. This phenomenon known as *rapid mixing* happens for a number of interesting examples. Section 5.4 presents a crucial tool used to show rapid mixing.

We used $\mathbf{x} \in \mathbf{R}^d$ to emphasize that distributions are multi-variate. From a Markov chain perspective, each value \mathbf{x} can take on is a state, i.e., a vertex of the graph on which the random walk takes place. Henceforth, we will use the subscripts i, j, k, \ldots to denote states and will use p_i instead of $p(x_1, x_2, \ldots, x_d)$ to denote the probability of the state corresponding to a given set of values for the variables. Recall that in the Markov chain terminology, vertices of the graph are called states.

Recall the notation that \mathbf{p}_t is the row vector of probabilities of the random walk being at each state (vertex of the graph) at time t. So, \mathbf{p}_t has as many components as there are states and its i^{th} component, p_{ti} , is the probability of being in state i at time t. Recall the long-term t-step average is

$$\mathbf{a}_{\mathbf{t}} = \frac{1}{t} \left[\mathbf{p}_{\mathbf{0}} + \mathbf{p}_{\mathbf{1}} + \dots + \mathbf{p}_{\mathbf{t}-\mathbf{1}} \right].$$
 (5.1)

The expected value of the function f under the probability distribution \mathbf{p} is $E(f) = \sum_i f_i p_i$ where f_i is the value of f at state i. Our estimate of this quantity will be the average value of f at the states seen in a t step walk. Call this estimate γ . Clearly, the expected value of γ is

$$E(\gamma) = \sum_{i} f_i\left(\frac{1}{t}\sum_{j=1}^{t} \operatorname{Prob}\left(\text{walk is in state } i \text{ at time } j\right)\right) = \sum_{i} f_i a_{ti}$$

The expectation here is with respect to the "coin tosses" of the algorithm, not with respect to the underlying distribution p. Let f_{max} denote the maximum absolute value of f. It is easy to see that

$$\left|\sum_{i} f_{i} p_{i} - E(\gamma)\right| \leq f_{\max} \sum_{i} |p_{i} - a_{ti}| = f_{\max} ||\mathbf{p} - \mathbf{a}_{t}||_{1}$$
(5.2)

where the quantity $||\mathbf{p} - \mathbf{a}_t||_1$ is the l_1 distance between the probability distributions \mathbf{p} and \mathbf{a}_t and is often called the "total variation distance" between the distributions. We will build tools to upper bound $||\mathbf{p} - \mathbf{a}_t||_1$. Since \mathbf{p} is the stationary distribution, the t for which $||\mathbf{p} - \mathbf{a}_t||_1$ becomes small is determined by the rate of convergence of the Markov chain to its steady state.

The following proposition is often useful.

Proposition 5.4 For two probability distributions **p** and **q**,

$$||\mathbf{p} - \mathbf{q}||_1 = 2\sum_i (p_i - q_i)^+ = 2\sum_i (q_i - p_i)^+$$

where $x^+ = x$ if $x \ge 0$ and $x^+ = 0$ if x < 0.

The proof is left as an exercise.

5.2.1 Metropolis-Hasting Algorithm

The Metropolis-Hasting algorithm is a general method to design a Markov chain whose stationary distribution is a given target distribution p. Start with a connected undirected graph G on the set of states. If the states are the lattice points (x_1, x_2, \ldots, x_d) in \mathbb{R}^d with $x_i \in \{0, 1, 2, \ldots, n\}$, then G could be the lattice graph with 2d coordinate edges at each interior vertex. In general, let r be the maximum degree of any vertex of G. The transitions of the Markov chain are defined as follows. At state i select neighbor j with probability $\frac{1}{r}$. Since the degree of i may be less than r, with some probability no edge is selected and the walk remains at i. If a neighbor j is selected and $p_j \ge p_i$, go to j. If $p_j < p_i$, go to j with probability p_j/p_i and stay at i with probability $1 - \frac{p_j}{p_i}$. Intuitively, this favors "heavier" states with higher p values. So, for i, adjacent to j in G,

$$p_{ij} = \frac{1}{r} \min\left(1, \frac{p_j}{p_i}\right)$$

and

$$p_{ii} = 1 - \sum_{j \neq i} p_{ij}.$$

Thus,

$$p_i p_{ij} = \frac{p_i}{r} \min\left(1, \frac{p_j}{p_i}\right) = \frac{1}{r} \min(p_i, p_j) = \frac{p_j}{r} \min\left(1, \frac{p_i}{p_j}\right) = p_j p_{ji}.$$

By Lemma 5.3, the stationary probabilities are indeed p_i as desired.

$$p(d) = p(a)p(a \to d) + p(c)p(c \to d) + p(d)p(d \to d)$$

= $\frac{1}{2} \frac{1}{12} + \frac{1}{8} \frac{1}{3} + \frac{1}{8} \frac{1}{3} = \frac{1}{8}$

Figure 5.2: Using the Metropolis-Hasting algorithm to set probabilities for a random walk so that the stationary probability will be the desired probability.

Example: Consider the graph in Figure 5.2. Using the Metropolis-Hasting algorithm, assign transition probabilities so that the stationary probability of a random walk is $p(a) = \frac{1}{2}$, $p(b) = \frac{1}{4}$, $p(c) = \frac{1}{8}$, and $p(d) = \frac{1}{8}$. The maximum degree of any vertex is three, so at *a*, the probability of taking the edge (a, b) is $\frac{1}{3}\frac{1}{4}\frac{2}{1}$ or $\frac{1}{6}$. The probability of taking the edge (a, c) is $\frac{1}{3}\frac{1}{8}\frac{2}{1}$ or $\frac{1}{12}$ and of taking the edge (a, d) is $\frac{1}{3}\frac{1}{8}\frac{2}{1}$ or $\frac{1}{12}$. Thus, the probability of staying at *a* is $\frac{2}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of taking the edge from *b* to *a* is $\frac{1}{3}$. The probability of a is $\frac{1}{4}\frac{1}{3} + \frac{1}{8}\frac{1}{3} + \frac{1}{2}\frac{1}{3} = \frac{1}{2}$, which is the desired probability.

5.2.2 Gibbs Sampling

Gibbs sampling is another Markov Chain Monte Carlo method to sample from a multivariate probability distribution. Let $p(\mathbf{x})$ be the target distribution where $\mathbf{x} = (x_1, \ldots, x_d)$. Gibbs sampling consists of a random walk on an undirectd graph whose vertices correspond to the values of $\mathbf{x} = (x_1, \ldots, x_d)$ and in which there is an edge from \mathbf{x} to \mathbf{y} if \mathbf{x} and \mathbf{y} differ in only one coordinate. Thus, the underlying graph is like a

d-dimensional lattice except that the vertices in the same coordinate line form a clique.

To generate samples of $\mathbf{x} = (x_1, \ldots, x_d)$ with a target distribution $p(\mathbf{x})$, the Gibbs sampling algorithm repeats the following steps. One of the variables x_i is chosen to be updated. Its new value is chosen based on the marginal probability of x_i with the other variables fixed. There are two commonly used schemes to determine which x_i to update. One scheme is to choose x_i randomly, the other is to choose x_i by sequentially scanning from x_1 to x_d .

Suppose that \mathbf{x} and \mathbf{y} are two states that differ in only one coordinate. Without loss of generality let that coordinate be the first. Then, in the scheme where a coordinate is randomly chosen to modify, the probability $p_{\mathbf{xy}}$ of going from \mathbf{x} to \mathbf{y} is

$$p_{\mathbf{x}\mathbf{y}} = \frac{1}{d}p(y_1|x_2, x_3, \dots, x_d).$$

The normalizing constant is 1/d since $\sum_{y_1} p(y_1|x_2, x_3, \ldots, x_d)$ equals 1 and summing over d coordinates gives a value of d. Similarly,

$$p_{\mathbf{yx}} = \frac{1}{d} p(x_1 | y_2, y_3, \dots, y_d)$$

= $\frac{1}{d} p(x_1 | x_2, x_3, \dots, x_d).$

Here use was made of the fact that for $j \neq 1$, $x_j = y_j$.

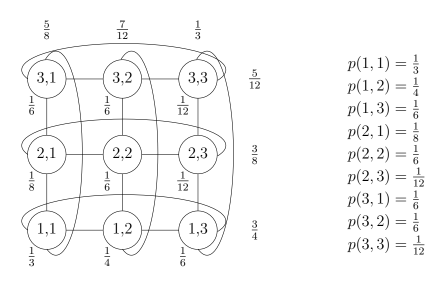
It is simple to see that this chain has stationary probability proportional to $p(\mathbf{x})$. Rewrite $p_{\mathbf{xy}}$ as

$$p_{\mathbf{x}\mathbf{y}} = \frac{1}{d} \frac{p(y_1 | x_2, x_3, \dots, x_d) p(x_2, x_3, \dots, x_d)}{p(x_2, x_3, \dots, x_d)}$$
$$= \frac{1}{d} \frac{p(y_1, x_2, x_3, \dots, x_d)}{p(x_2, x_3, \dots, x_d)}$$
$$= \frac{1}{d} \frac{p(\mathbf{y})}{p(x_2, x_3, \dots, x_d)}$$

again using $x_j = y_j$ for $j \neq 1$. Similarly write

$$p_{\mathbf{yx}} = \frac{1}{d} \frac{p(\mathbf{x})}{p(x_2, x_3, \dots, x_d)}$$

from which it follows that $p(\mathbf{x})p_{xy} = p(\mathbf{y})p_{yx}$. By Lemma 5.3 the stationary probability of the random walk is $p(\mathbf{x})$.



 $p_{(11)(12)} = \frac{1}{d} p_{12} / (p_{11} + p_{12} + p_{13} = \frac{1}{2} \frac{1}{4} / (\frac{1}{3} \frac{1}{4} \frac{1}{6} = \frac{1}{2} \frac{1}{4} / \frac{9}{12} = \frac{1}{2} \frac{1}{4} \frac{4}{3} = \frac{1}{6}$ Calculation of edge probability $p_{(11)(12)}$

$p_{(11)(12)} = \frac{1}{2}\frac{1}{4}\frac{4}{3} = \frac{1}{6}$	$p_{(12)(11)} = \frac{1}{2} \frac{1}{3} \frac{1}{3} = \frac{2}{9}$	$p_{(13)(11)} = \frac{1}{2} \frac{1}{3} \frac{4}{3} = \frac{2}{9}$	$p_{(21)(22)} = \frac{1}{2} \frac{1}{6} \frac{1}{8} = \frac{2}{9}$
$p_{(11)(13)} = \frac{1}{2}\frac{1}{6}\frac{4}{3} = \frac{1}{9}$	$p_{(12)(13)} = \frac{1}{2} \frac{1}{6} \frac{1}{3} = \frac{1}{9}$	$p_{(13)(12)} = \frac{1}{2}\frac{1}{4}\frac{4}{3} = \frac{1}{6}$	$p_{(21)(23)} = \frac{1}{2} \frac{1}{12} \frac{8}{3} = \frac{1}{9}$
$p_{(11)(21)} = \frac{1}{2} \frac{1}{8} \frac{8}{5} = \frac{1}{10}$	$p_{(12)(22)} = \frac{1}{2} \frac{1}{6} \frac{12}{7} = \frac{1}{7}$	$p_{(13)(23)} = \frac{1}{2} \frac{1}{12} \frac{3}{1} = \frac{1}{8}$	$p_{(21)(11)} = \frac{1}{2} \frac{1}{3} \frac{8}{5} = \frac{4}{15}$
$p_{(11)(31)} = \frac{1}{2} \frac{1}{6} \frac{8}{5} = \frac{2}{15}$	$p_{(12)(32)} = \frac{1}{2} \frac{1}{6} \frac{12}{7} = \frac{1}{7}$	$p_{(13)(33)} = \frac{1}{2} \frac{1}{12} \frac{3}{1} = \frac{1}{8}$	$p_{(21)(31)} = \frac{1}{2} \frac{1}{6} \frac{1}{5} = \frac{2}{15}$
Edge probabilities.			

 $p_{11}p_{(11)(12)} = \frac{1}{3}\frac{1}{6} = \frac{1}{4}\frac{2}{9} = p_{12}p_{(12)(11)}$ $p_{11}p_{(11)(13)} = \frac{1}{3}\frac{1}{9} = \frac{1}{6}\frac{2}{9} = p_{13}p_{(13)(11)}$ $p_{11}p_{(11)(21)} = \frac{1}{3}\frac{1}{10} = \frac{1}{8}\frac{4}{15} = p_{21}p_{(21)(11)}$ Verification of a few edges.

Note that the edge probabilities out of a state such as (1,1) do not add up to one. That is, with some probability the walk stays at the state that it is in. For example, $p_{(11)(11)} = 1 - (p_{(11)(12)} + p_{(11)(13)} + p_{(11)(21)} + p_{(11)(31)}) = 1 - \frac{1}{6} - \frac{1}{24} - \frac{1}{32} - \frac{1}{24} = \frac{9}{32}$.

Figure 5.3: Using the Gibbs algorithm to set probabilities for a random walk so that the stationary probability will be a desired probability.

5.3 Areas and Volumes

Computing areas and volumes is a classical problem. For many regular figures in two and three dimensions there are closed form formulae. In Chapter 2, we saw how to compute volume of a high dimensional sphere by integration. For general convex sets in d-space, there are no closed form formulae. Can we estimate volumes of d-dimensional convex sets in time that grows as a polynomial function of d? The MCMC method answes this question in the affirmative.

One way to estimate the area of the region is to enclose it in a rectangle and estimate the ratio of the area of the region to the area of the rectangle by picking random points in the rectangle and seeing what proportion land in the region. Such methods fail in high dimensions. Even for a sphere in high dimension, a cube enclosing the sphere has exponentially larger area, so exponentially many samples are required to estimate the volume of the sphere.

It turns out that the problem of estimating volumes of sets is reducible to the problem of drawing uniform random samples from sets. Suppose one wants to estimate the volume of a convex set R. Create a concentric series of larger and larger spheres S_1, S_2, \ldots, S_k such that S_1 is contained in R and S_k contains R. Then

$$\operatorname{Vol}(R) = \operatorname{Vol}(S_k \cap R) = \frac{\operatorname{Vol}(S_k \cap R)}{\operatorname{Vol}(S_{k-1} \cap R)} \frac{\operatorname{Vol}(S_{k-1} \cap R)}{\operatorname{Vol}(S_{k-2} \cap R)} \cdots \frac{\operatorname{Vol}(S_2 \cap R)}{\operatorname{Vol}(S_1 \cap R)} \operatorname{Vol}(S_1)$$

If the radius of the sphere S_i is $1 + \frac{1}{d}$ times the radius of the sphere S_{i-1} , then the value of

$$\frac{\operatorname{Vol}(S_{k-1}\cap R)}{\operatorname{Vol}(S_{k-2}\cap R)}$$

can be estimated by rejection sampling provided one can select points at random from a d-dimensional region. Since the radii of the spheres grows as $1 + \frac{1}{d}$, the volume of a sphere is $\left(1 + \frac{1}{d}\right)^d < e$ times the volume of the preceding sphere and the number of spheres is at most

$$O(\log_{1+(1/d)} r) = O(rd)$$

where r is the ratio of the radius of S_k to the radius of S_1 .

It remains to show how to draw a uniform random sample from a *d*-dimensional set. It is at this point that we require the set to be convex so that the Markov chain technique will converge quickly to its stationary probability. To select a random sample from a *d*-dimensional convex set, impose a grid on the region and do a random walk on the grid points. At each time, pick one of the 2*d* coordinate neighbors of the current grid point, each with probability 1/(2d) and go to the neighbor if it is still in the set; otherwise, stay put and repeat. If the grid length in each of the *d* coordinate directions is at most some *a*, the total number of grid points in the set is at most a^d . Although this is exponential in *d*, the Markov chain turns out to be rapidly mixing (the proof is beyond our scope here)

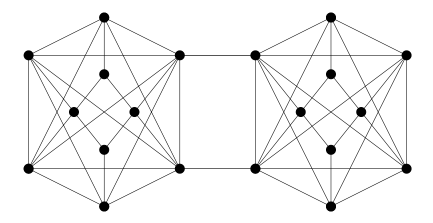


Figure 5.4: A network with a constriction.

and leads to polynomial time bounded algorithm to estimate the volume of any convex set in \mathbf{R}^d .

5.4 Convergence of Random Walks on Undirected Graphs

The Metropolis-Hasting algorithm and Gibbs sampling both involve a random walk. Initial states of the walk are highly dependent on the start state of the walk. Both these walks are random walks on edge-weighted undirected graphs. Such Markov chains are derived from electrical networks. Recall the following notation which we will use throughout this section. Given a network of resistors, the conductance of edge (x, y)is denoted c_{xy} and the normalizing constant c_x equals $\sum_y c_{xy}$. The Markov chain has transition probabilities $p_{xy} = c_{xy}/c_x$. We assume the chain is connected. Since

$$c_x p_{xy} = c_{xy} = c_{yx} = c_y c_{yx} / c_y = c_y p_{xy}$$

the stationary probabilities are proportional to c_x where the normalization constant is $c_0 = \sum_x c_x$.

An important question is how fast the walk starts to reflect the stationary probability of the Markov process. If the convergence time was proportional to the number of states, the algorithms would not be very useful since the number of states can be exponentially large.

There are clear examples of connected chains that take a long time to converge. A chain with a constriction, see Figure 5.4, takes a long time to converge since the walk is unlikely to reach the narrow passage between the two halves, both of which are reasonably big. We will show in Theorem 5.5 that the time to converge is quantitatively related to

the tightest constriction.

A function is unimodal if it has a single maximum, i.e., it increases and then decreases. A unimodal function like the normal density has no constriction blocking a random walk from getting out of a large set of states, whereas a bimodal function can have a constriction. Interestingly, many common multivariate distributions as well as univariate probability distributions like the normal and exponential are unimodal and sampling according to these distributions can be done using the methods here.

A natural problem is estimating the probability of a convex region in d-space according to a normal distribution. One technique to do this is rejection sampling. Let R be the region defined by the inequality $x_1 + x_2 + \cdots + x_{d/2} \leq x_{d/2+1} + \cdots + x_d$. Pick a sample according to the normal distribution and accept the sample if it satisfies the inequality. If not, reject the sample and retry until one gets a number of samples satisfying the inequality. The probability of the region is approximated by the fraction of the samples that satisfied the inequality. However, suppose R was the region $x_1 + x_2 + \cdots + x_{d-1} \leq x_d$. The probability of this region is exponentially small in d and so rejection sampling runs into the problem that we need to pick exponentially many samples before we accept even one sample. This second situation is typical. Imagine computing the probability of failure of a system. The object of design is to make the system reliable, so the failure probability is likely to be very low and rejection sampling will take a long time to estimate the failure probability.

In general, there could be constrictions that prevent rapid convergence of a Markov chain to its stationary probability. However, if the set is convex in any number of dimensions, then there are no constrictions and there is rapid convergence although the proof of this is beyond the scope of this book.

Recall the notation that π stands for the stationary probability distribution of a connected Markov Chain. We define below a combinatorial measure of constriction for a Markov chain, called the *normalized conductance*. We will relate normalized conductance to the time by which the average probability distribution of the chain is guaranteed to be close to the stationarity probability distribution. We call this ε -mixing time:

Definition 5.1 Fix $\varepsilon > 0$. The ε -mixing time of a Markov chain is the minimum integer t such that for any starting distribution \mathbf{p}_0 , the 1-norm distance between the t-step running average probability distribution 16 and the stationary distribution is at most ε .

Definition 5.2 For a subset S of vertices, let $\pi(S)$ denote $\sum_{x \in S} \pi_x$. The normalized conductance $\Phi(S)$ of S is

$$\Phi(S) = \frac{\sum_{(x,y)\in(S,\bar{S})} \pi_x p_{xy}}{\min(\pi(S), \pi(\bar{S}))}.$$

¹⁶Recall that $\mathbf{a}_{\mathbf{t}} = \frac{1}{t}(\mathbf{p}_0 + \mathbf{p}_1 + \dots + \mathbf{p}_{t-1})$ is called the running average distribution.

There is a simple interpretation of $\Phi(S)$. Suppose without loss of generality that $\pi(S) \leq \pi(\bar{S})$. Then, we may write $\Phi(S)$ as

$$\phi(S) = \sum_{x \in S} \frac{\pi_x}{\underbrace{\pi(S)}_a} \underbrace{\sum_{y \in \bar{S}} p_{xy}}_{b}.$$

Here, a is the probability of being in x if we were in the stationary distribution restricted to S and b is the probability of stepping from x to \bar{S} in a single step. It is easy to show that if we started in the distribution $p_{0,x} = \pi_s/\pi(S)$ for $x \in S$ (and 0 on \bar{S}), the expected number of steps before we step into \bar{S} is

$$1\Phi(S) + 2(1 - \Phi(S))\Phi(S) + 3(1 - \Phi(S))^2\Phi(S) + \dots = \frac{1}{\Phi(S)}.$$

Clearly, to be close to the stationary distribution, we must at least get to \overline{S} once. So, mixing time is lower bounded by $1/\Phi(S)$. Since we could have taken any S, mixing time is lower bounded by the minimum over all S of $\Phi(S)$. We define this quantity to be the normalized conductance of the Markov Chain:

Definition 5.3 The normalized conductance of the Markov chain, denoted Φ , is defined by

$$\Phi = \min_{S} \Phi(S).$$

As we just argued, normalized conductance being high is a necessary condition for mixing. The theorem below proves the converse: normalized conductance being high is sufficient for mixing. Intuitively, if Φ is large, the walk rapidly leaves any subset of states. But the proof of the theorem is quite difficult. After we prove it, we will see examples where the mixing time is much smaller than the cover time. That is, the number of steps before a random walk reaches a random state independent of its starting state is much smaller than the average number of steps needed to reach every state. In fact for some graphs, called expenders, the mixing time is logarithmic in the number of states.

Theorem 5.5 The ε -mixing time of a random walk on an undirected graph is

$$O\left(\frac{\ln(1/\pi_{min})}{\Phi^2\varepsilon^3}\right)$$

where π_{min} is the minimum stationary probability of any state.

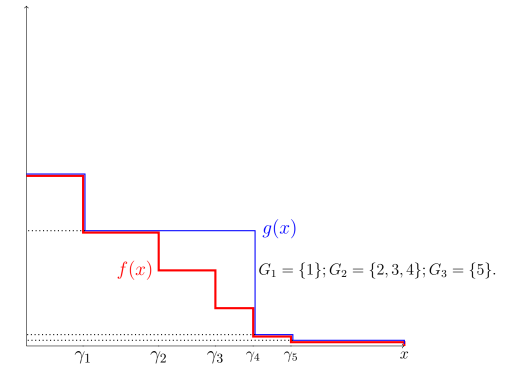


Figure 5.5: Bounding l_1 distance.

Proof: Let $t = \frac{c \ln(1/\pi_{\min})}{\Phi^2 \varepsilon^3}$, for a suitable constant c. Let

$$\mathbf{a} = \mathbf{a}_{\mathbf{t}} = \frac{1}{t}(\mathbf{p}_0 + \mathbf{p}_1 + \dots + \mathbf{p}_{t-1})$$

be the running average distribution. We need to show that $||\mathbf{a} - \boldsymbol{\pi}||_1 \leq \varepsilon$. Let

$$v_i = \frac{a_i}{\pi_i}.$$

The proof has two parts - the first is technical reducing what we need to prove by a series of manipulations to proving essentially that v_i do not drop too fast as we increase *i*. [Note: In the extreme, if all v_i equal 1, $||\mathbf{a} - \boldsymbol{\pi}||_1 = 0$; so, intuitively, indeed, proving that v_i do not fall too much should give us what we want.] In the second part, we prove that v_i do not fall fast using the concept of "probability flows".

Renumber states so that $v_1 \ge v_2 \ge \cdots$. We call a state *i* for which $v_i > 1$ "heavy" since it has more probability according to **a** than its stationary probability. Let i_0 be the maximum *i* such that $v_i > 1$; it is the last heavy state. By Proposition (5.4):

$$||\mathbf{a} - \boldsymbol{\pi}||_{1} = 2\sum_{i=1}^{i_{0}} (v_{i} - 1)\pi_{i} = 2\sum_{i \ge i_{0} + 1} (1 - v_{i})\pi_{i}.$$
(5.3)

Let

$$\gamma_i = \pi_1 + \pi_2 + \dots + \pi_i.$$

Define a function $f : [0, \gamma_{i_0}] \rightarrow \text{Reals by } f(x) = v_i - 1 \text{ for } x \in [\gamma_{i-1}, \gamma_i)$. See figure (5.5). Now,

$$\sum_{i=1}^{i_0} (v_i - 1)\pi_i = \int_0^{\gamma_{i_0}} f(x) \, dx.$$
(5.4)

We make one more technical modification. We divide $\{1, 2, \ldots, i_0\}$ into groups $G_1, G_2, G_3, \ldots, G_r$, of contiguous subsets. [We specify the groups later.] Let $u_t = \operatorname{Max}_{i \in G_t} v_i$. Then we define a new function g(x) by $g(x) = u_t$ by (see figure (5.5)): for $x \in \bigcup_{i \in G_t} [\gamma_{i-1}, \gamma_i)$. Clearly,

$$\int_{0}^{\gamma_{i_{0}}} f(x) \, dx \le \int_{0}^{\gamma_{i_{0}}} g(x) \, dx. \tag{5.5}$$

We now assert (with $u_{r+1} = 0$):

$$\int_0^{\gamma_{i_0}} g(x) \, dx = \sum_{t=1}^r \pi(G_1 \cup G_2 \cup \ldots \cup G_t)(u_t - u_{t+1}).$$
(5.6)

This is just the statement that the area under g(x) in the figure is exactly covered by the rectangles whose bottom sides are the dotted lines. We leave the formal proof of this to the reader. We now focus on proving that

$$\sum_{t=1}^{r} \pi(G_1 \cup G_2 \cup \ldots \cup G_t)(u_t - u_{t+1}) \le \varepsilon/2,$$
(5.7)

(for a sub-division into groups we specify) which suffices by (5.3, 5.4, 5.5 and 5.6). While we start the proof of (5.7) with a technical observation (5.8), its proof will involve two nice ideas: the notion of probability flow and reckoning probability flow in two different ways. First, the technical observation: if $2\sum_{i\geq i_0+1}(1-v_i)\pi_i \leq \varepsilon$ and we would be done by (5.3). So assume now that $\sum_{i\geq i_0+1}(1-v_i)\pi_i \geq \varepsilon/2$ from which it follows that $\sum_{i\geq i_0+1}\pi_i \geq \varepsilon/2$ and so, for any subset A of heavy nodes,

$$\operatorname{Min}(\pi(A), \pi(\bar{A})) \ge \frac{\varepsilon}{2}\pi(A).$$
(5.8)

We now define the subsets. G_1 will be just $\{1\}$. In general, suppose $G_1, G_2, \ldots, G_{t-1}$ have already been defined. We start G_t at $i_t = 1+$ (end of G_{t-1}). Let $i_t = k$. We will define l, the last element of G_t to be the largest integer greater than or equal to k and at most i_0 so that

$$\sum_{j=k+1}^{l} \pi_j \le \varepsilon \Phi \gamma_k / 4.$$

Lemma 5.6 Suppose groups $G_1, G_2, \ldots, G_r, u_1.u_2, \ldots, u_r, u_{r+1}$ are as above. Then,

$$\pi(G_1 \cup G_2 \cup \ldots G_r)(u_t - u_{t+1}) \le \frac{8}{t\Phi\varepsilon}.$$

Proof: This is the main Lemma. The proof of the Lemma uses a crucial idea of probability flows. We will use two ways of calculating the probability flow from heavy states to ligh states when we execute one step of the Markov chain starting at probabilities **a**. The probability vector after that step is $\mathbf{a}P$. Now, $\mathbf{a} - \mathbf{a}P$ is the net loss of probability for each state due to the step.

Consider a particular group $G_t = \{k, k+1, \ldots, l\}$, say. First consider the case when $k < i_0$. Let $A = \{1, 2, \ldots, k\}$. The net loss of probability for each state from the set A in one step is $\sum_{i=1}^{k} (a_i - (\mathbf{a}P)_i)$ which is at most $\frac{2}{t}$ by the proof of Theorem 5.2.

Another way to reckon the net loss of probability from A is to take the difference of the probability flow from A to \overline{A} and the flow from \overline{A} to A. For any i < j,

net-flow
$$(i, j) =$$
flow $(i, j) -$ flow $(j, i) = \pi_i p_{ij} v_i - \pi_j p_{ji} v_j = \pi_j p_{ji} (v_i - v_j) \ge 0,$

Thus, for any two states i, j, with i heavier than j, i.e., i < j, there is a non-negative net flow from i to j. [This is intuitvely reasonable, since, it says that probability is flowing from heavy to light states.] Since, $l \ge k$, the flow from A to $\{k + 1, k + 2, \ldots, l\}$ minus the flow from $\{k + 1, k + 2, \ldots, l\}$ to A is nonnegative. Since for $i \le k$ and j > l, we have $v_i \ge v_k$ and $v_j \le v_{l+1}$, the net loss from A is at least

$$\sum_{\substack{i \le k \\ j > l}} \pi_j p_{ji} (v_i - v_j) \ge (v_k - v_{l+1}) \sum_{\substack{i \le k \\ j > l}} \pi_j p_{ji}.$$

Thus,

$$(v_k - v_{l+1}) \sum_{\substack{i \le k \\ j > l}} \pi_j p_{ji} \le \frac{2}{t}.$$
 (5.9)

Since

$$\sum_{i=1}^{k} \sum_{j=k+1}^{l} \pi_j p_{ji} \le \sum_{j=k+1}^{l} \pi_j \le \varepsilon \Phi \pi(A)/4$$

and by the definition of Φ , using (5.8)

$$\sum_{\leq k < j} \pi_j p_{ji} \ge \Phi \operatorname{Min}(\pi(A), \pi(\bar{A})) \ge \varepsilon \Phi \gamma_k / 2,$$

we have, $\sum_{\substack{i \leq k \\ j > l}} \pi_j p_{ji} = \sum_{i \leq k < j} \pi_j p_{ji} - \sum_{i \leq k; j \leq l} \pi_j p_{ji} \geq \varepsilon \Phi \gamma_k / 4$. Substituting this into the

inequality (5.9) gives

$$v_k - v_{l+1} \le \frac{8}{t\varepsilon\Phi\gamma_k},\tag{5.10}$$

proving the Lemma provided $k < i_0$. If $k = i_0$, the proof is similar (but simpler).

Now to prove (5.7), we now only need an upper bound on r, the number of groups. If $G_t = \{k, k+1, \ldots, l\}$, with $l < i_0$, then by definition of l, we have $\gamma_{l+1} \ge (1 + \frac{\varepsilon \Phi}{2})\gamma_k$. So, $r \le \ln_{1+(\varepsilon \Phi/2)}(1/\pi_1) + 2 \le \ln(1/\pi_1)/(\varepsilon \Phi/2) + 2$. This completes the proof of (5.7) and the theorem.

5.4.1 Using Normalized Conductance to Prove Convergence

We now apply Theorem 5.5 to some examples to illustrate how the normalized conductance bounds the rate of convergence. Our first examples will be simple graphs. The graphs do not have rapid converge, but their simplicity helps illustrate how to bound the normalized conductance and hence the rate of convergence.

A 1-dimensional lattice

Consider a random walk on an undirected graph consisting of an *n*-vertex path with self-loops at the both ends. With the self loops, the stationary probability is a uniform $\frac{1}{n}$ over all vertices. The set with minimum normalized conductance is the set with probability $\pi \leq \frac{1}{2}$ and the maximum number of vertices with the minimum number of edges leaving it. This set consists of the first n/2 vertices, for which total conductance of edges from S to \bar{S} is (with m = n/2) $\pi_m p_{m,m+1} = \Omega(\frac{1}{n})$ and $\pi(S) = \frac{1}{2}$. (π_m is the stationary probability of vertex numbered m.) Thus

$$\Phi(S) = 2\pi_m \ p_{m,m+1} = \Omega(1/n).$$

By Theorem 5.5, for ε a constant such as 1/100, after $O(n^2 \log n)$ steps, $||\mathbf{a}_t - \boldsymbol{\pi}||_1 \leq 1/100$. This graph does not have rapid convergence. The hitting time and the cover time are $O(n^2)$. In many interesting cases, the mixing time may be much smaller than the cover time. We will see such an example later.

A 2-dimensional lattice

Consider the $n \times n$ lattice in the plane where from each point there is a transition to each of the coordinate neighbors with probability 1/4. At the boundary there are self-loops with probability 1-(number of neighbors)/4. It is easy to see that the chain is connected. Since $p_{ij} = p_{ji}$, the function $f_i = 1/n^2$ satisfies $f_i p_{ij} = f_j p_{ji}$ and by Lemma 5.3 is the stationary probability. Consider any subset S consisting of at most half the states. Index states by their x and y coordinates. For at least half the states in S, either row x or column y intersects \overline{S} (Exercise 5.4). So at least $\Omega(|S|/n)$ points in S are adjacent to points in \overline{S} . Each such point contributes $\pi_i p_{ij} = \Omega(1/n^2)$ to flow (S, \overline{S}) . So

$$\sum_{i \in S} \sum_{j \in \bar{S}} \pi_i p_{ij} \ge c|S|/n^3$$

Thus, $\Phi \ge \Omega(1/n)$. By Theorem 5.5, after $O(n^2 \ln n/\varepsilon^2)$ steps, $|\mathbf{a}_t - \boldsymbol{\pi}|_1 \le 1/100$.

A lattice in *d*-dimensions

Next consider the $n \times n \times \cdots \times n$ lattice in *d*-dimensions with a self-loop at each boundary point with probability 1 - (number of neighbors)/2d. The self loops make all π_i equal to n^{-d} . View the lattice as an undirected graph and consider the random walk

on this undirected graph. Since there are n^d states, the cover time is at least n^d and thus exponentially dependent on d. It is possible to show (Exercise 5.20) that Φ is $\Omega(1/dn)$. Since all π_i are equal to n^{-d} , the mixing time is $O(d^3n^2 \ln n/\varepsilon^2)$, which is polynomially bounded in n and d.

The *d*-dimensional lattice is related to the Metropolis-Hastings algorithm and Gibbs sampling although in those constructions there is a nonuniform probability distribution at the vertices. However, the *d*-dimension lattice case suggests why the Metropolis-Hastings and Gibbs sampling constructions might converge fast.

A clique

Consider an *n* vertex clique with a self loop at each vertex. For each edge, $p_{xy} = \frac{1}{n}$ and thus for each vertex, $\pi_x = \frac{1}{n}$. Let *S* be a subset of the vertices. Then

$$\sum_{x \in S} \pi_x = \frac{|S|}{n}.$$
$$\sum_{(x,y) \in (S,\bar{S})} \pi_x p_{xy} = \frac{1}{n^2} |S| |\overline{S}|$$

and

$$\Phi(S) = \frac{\sum_{(x,y)\in(S,\bar{S})} \pi_x p_{xy}}{\sum_{x\in S} \pi_x \sum_{x\in\bar{S}} \pi_x} = 1.$$

This gives a ε -mixing time of

$$O\left(\frac{\ln\frac{1}{\pi_{\min}}}{\Phi^2\epsilon^3}\right) = O\left(\frac{\ln n}{\epsilon^3}\right).$$

A connected undirected graph

Next consider a random walk on a connected n vertex undirected graph where at each vertex all edges are equally likely. The stationary probability of a vertex equals the degree of the vertex divided by the sum of degrees. The sum of the vertex degrees is at most n^2 and thus, the steady state probability of each vertex is at least $\frac{1}{n^2}$. Since the degree of a vertex is at most n, the probability of each edge at a vertex is at least $\frac{1}{n}$. For any S, the total conductance of edges out of S is greater than or equal to

$$\frac{1}{n^2}\frac{1}{n} = \frac{1}{n^3}.$$

Thus, Φ is at least $\frac{1}{n^3}$. Since $\pi_{\min} \geq \frac{1}{n^2}$, $\ln \frac{1}{\pi_{\min}} = O(\ln n)$. Thus, the mixing time is $O(n^6 \ln n/\varepsilon^3)$.

The Gaussian distribution on the interval [-1,1]

Consider the interval [-1, 1]. Let δ be a "grid size" specified later and let G be the graph consisting of a path on the $\frac{2}{\delta} + 1$ vertices $\{-1, -1 + \delta, -1 + 2\delta, \dots, 1 - \delta, 1\}$ having self loops at the two ends. Let $\pi_x = ce^{-\alpha x^2}$ for $x \in \{-1, -1 + \delta, -1 + 2\delta, \dots, 1 - \delta, 1\}$ where $\alpha > 1$ and c has been adjusted so that $\sum_x \pi_x = 1$.

We now describe a simple Markov chain with the π_x as its stationary probability and argue its fast convergence. With the Metropolis-Hastings' construction, the transition probabilities are

$$p_{x,x+\delta} = \frac{1}{2} \min\left(1, \frac{e^{-\alpha(x+\delta)^2}}{e^{-\alpha x^2}}\right)$$
 and $p_{x,x-\delta} = \frac{1}{2} \min\left(1, \frac{e^{-\alpha(x-\delta)^2}}{e^{-\alpha x^2}}\right)$.

Let S be any subset of states with $\pi(S) \leq \frac{1}{2}$. First consider the case when S is an interval $[k\delta, 1]$ for $k \geq 2$. It is easy to see that

$$\pi(S) \leq \int_{x=(k-1)\delta}^{\infty} ce^{-\alpha x^2} dx$$
$$\leq \int_{(k-1)\delta}^{\infty} \frac{x}{(k-1)\delta} ce^{-\alpha x^2} dx$$
$$= O\left(\frac{ce^{-\alpha((k-1)\delta)^2}}{\alpha(k-1)\delta}\right).$$

Now there is only one edge from S to \overline{S} and total conductance of edges out of S is

$$\sum_{i\in S}\sum_{j\notin S}\pi_i p_{ij} = \pi_{k\delta} p_{k\delta,(k-1)\delta} = \min(ce^{-\alpha k^2\delta^2}, ce^{-\alpha(k-1)^2\delta^2}) = ce^{-\alpha k^2\delta^2}$$

Using $2 \le k \le 1/\delta$, $\alpha \ge 1$, and $\pi(\bar{S}) \le 1$, we have

$$\Phi(S) = \frac{\text{flow}(S, \bar{S})}{\pi(S)\pi(\bar{S})} \ge ce^{-\alpha k^2 \delta^2} \frac{\alpha(k-1)\delta}{ce^{-\alpha((k-1)\delta)^2}}$$
$$\ge \Omega(\alpha(k-1)\delta e^{-\alpha \delta^2(2k-1)}) \ge \Omega(\alpha \delta e^{-O(\alpha \delta)})$$

For $\delta < \frac{1}{\alpha}$, we have $\alpha \delta < 1$, so $e^{-O(\alpha \delta)} = \Omega(1)$, thus, $\Phi(S) \ge \Omega(\alpha \delta)$. Now, $\pi_{\min} \ge ce^{-\alpha} \ge e^{-1/\delta}$, so $\ln(1/\pi_{\min}) \le 1/\delta$.

If S is not an interval of the form [k, 1] or [-1, k], then the situation is only better since there is more than one "boundary" point which contributes to flow (S, \overline{S}) . We do not present this argument here. By Theorem 5.5 in $\Omega(1/\alpha^2\delta^3\varepsilon^3)$ steps, a walk gets within ε of the steady state distribution.

In these examples, we have chosen simple probability distributions. The methods extend to more complex situations.

5.5 Electrical Networks and Random Walks

In the next few sections, we study the relationship between electrical networks and random walks on undirected graphs. The graphs have nonnegative weights on each edge. A step is executed by picking a random edge from the current vertex with probability proportional to the edge's weight and traversing the edge.

An electrical network is a connected, undirected graph in which each edge (x, y) has a resistance $r_{xy} > 0$. In what follows, it is easier to deal with conductance defined as the reciprocal of resistance, $c_{xy} = \frac{1}{r_{xy}}$, rather than resistance. Associated with an electrical network is a random walk on the underlying graph defined by assigning a probability $p_{xy} = \frac{c_{xy}}{c_x}$ to the edge (x, y) incident to the vertex x, where the normalizing constant c_x equals $\sum_{y} c_{xy}$. Note that although c_{xy} equals c_{yx} , the probabilities p_{xy} and p_{yx} may not be equal due to the normalization required to make the probabilities at each vertex sum to one. We shall soon see that there is a relationship between current flowing in an electrical network and a random walk on the underlying graph.

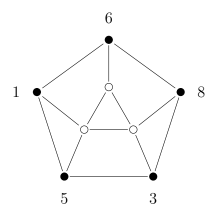
Since we assume that the undirected graph is connected, by Theorem 5.2 there is a unique stationary probability distribution. The stationary probability distribution is π where $\pi_x = \frac{c_x}{c_0}$ where $c_0 = \sum_x c_x$. To see this, for all x and y

$$\pi_x p_{xy} = \frac{c_x}{c_0} \frac{c_{xy}}{c_x} = \frac{c_{xy}}{c_0} = \frac{c_y}{c_0} \frac{c_{yx}}{c_y} = \pi_y p_{yx}$$

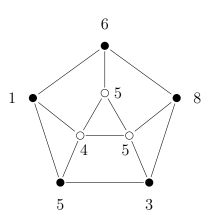
and hence by Lemma 5.3, π is the unique stationary probability.

Harmonic functions

Harmonic functions are useful in developing the relationship between electrical networks and random walks on undirected graphs. Given an undirected graph, designate a nonempty set of vertices as boundary vertices and the remaining vertices as interior vertices. A harmonic function g on the vertices is one in which the value of the function at the boundary vertices is fixed to some boundary condition and the value of g at any interior vertex x is a weighted average of the values at all the adjacent vertices y, with weights p_{xy} satisfying $\sum_{y} p_{xy} = 1$ for each x. Thus, if at every interior vertex x for some set of weights p_{xy} satisfying $\sum_{y} p_{xy} = 1$, $g_x = \sum_{y} g_y p_{xy}$, then g is an harmonic function.



Graph with boundary vertices dark and boundary conditions specified.



Values of harmonic function satisfying boundary conditions where the edge weights at each vertex are equal

Figure 5.6: Graph illustrating an harmonic function.

Example: Convert an electrical network with conductances c_{xy} to a weighted, undirected graph with probabilities p_{xy} . Let **f** be a function satisfying $\mathbf{f}P = \mathbf{f}$ where P is the matrix of probabilities. It follows that the function $g_x = \frac{f_x}{c_x}$ is harmonic.

$$g_x = \frac{f_x}{c_x} = \frac{1}{c_x} \sum_y f_y p_{yx} = \frac{1}{c_x} \sum_y f_y \frac{c_{yx}}{c_y}$$
$$= \frac{1}{c_x} \sum_y f_y \frac{c_{xy}}{c_y} = \sum_y \frac{f_y}{c_y} \frac{c_{xy}}{c_x} = \sum_y g_y p_{xy}$$

A harmonic function on a connected graph takes on its maximum and minimum on the boundary. Suppose the maximum does not occur on the boundary. Let S be the set of interior vertices at which the maximum value is attained. Since S contains no boundary vertices, \overline{S} is nonempty. Connectedness implies that there is at least one edge (x, y) with $x \in S$ and $y \in \overline{S}$. The value of the function at x is the weighted average of the value at its neighbors, all of which are less than or equal to the value at x and the value at y is strictly less, a contradiction. The proof for the minimum value is identical.

There is at most one harmonic function satisfying a given set of equations and boundary conditions. For suppose there were two solutions, f(x) and g(x). The difference of two solutions is itself harmonic. Since h(x) = f(x) - g(x) is harmonic and has value zero on the boundary, by the min and max principles it has value zero everywhere. Thus f(x) = g(x).

The analogy between electrical networks and random walks

There are important connections between electrical networks and random walks on undirected graphs. Choose two vertices a and b. For reference purposes let the voltage v_b equal zero. Attach a current source between a and b so that the voltage v_a equals one. Fixing the voltages at v_a and v_b induces voltages at all other vertices along with a current flow through the edges of the network. The analogy between electrical networks and random walks is the following. Having fixed the voltages at the vertices a and b, the voltage at an arbitrary vertex x equals the probability of a random walk starting at xreaching a before reaching b. If the voltage v_a is adjusted so that the current flowing into vertex a corresponds to one walk, then the current flowing through an edge is the net frequency with which a random walk from a to b traverses the edge.

Probabilistic interpretation of voltages

Before showing that the voltage at an arbitrary vertex x equals the probability of a random walk starting at x reaching a before reaching b, we first show that the voltages form a harmonic function. Let x and y be adjacent vertices and let i_{xy} be the current flowing through the edge from x to y. By Ohm's law,

$$i_{xy} = rac{v_x - v_y}{r_{xy}} = (v_x - v_y)c_{xy}.$$

By Kirchhoff's law the currents flowing out of each vertex sum to zero.

$$\sum_{y} i_{xy} = 0$$

Replacing currents in the above sum by the voltage difference times the conductance yields

$$\sum_{y} (v_x - v_y)c_{xy} = 0$$
$$v_x \sum_{y} c_{xy} = \sum_{y} v_y c_{xy}$$

or

Observing that
$$\sum_{y} c_{xy} = c_x$$
 and that $p_{xy} = \frac{c_{xy}}{c_x}$, yields $v_x c_x = \sum_{y} v_y p_{xy} c_x$. Hence, $v_x = \sum_{y} v_y p_{xy}$. Thus, the voltage at each vertex x is a weighted average of the voltages at the adjacent vertices. Hence the voltages form a harmonic function with $\{a, b\}$ as the boundary.

Let p_x be the probability that a random walk starting at vertex x reaches a before b. Clearly $p_a = 1$ and $p_b = 0$. Since $v_a = 1$ and $v_b = 0$, it follows that $p_a = v_a$ and $p_b = v_b$. Furthermore, the probability of the walk reaching a from x before reaching b is the sum over all y adjacent to x of the probability of the walk going from x to y in the first step and then reaching a from y before reaching b. That is

$$p_x = \sum_y p_{xy} p_y$$

Hence, p_x is the same harmonic function as the voltage function v_x and \mathbf{v} and \mathbf{p} satisfy the same boundary conditions at a and b. Thus, they are identical functions. The probability of a walk starting at x reaching a before reaching b is the voltage v_x .

Probabilistic interpretation of current

In a moment, we will set the current into the network at a to have a value which we will equate with one random walk. We will then show that the current i_{xy} is the net frequency with which a random walk from a to b goes through the edge xy before reaching b. Let u_x be the expected number of visits to vertex x on a walk from a to b before reaching b. Clearly $u_b = 0$. Every time the walk visits x, x not equal to a, it must come to x from some vertex y. Thus, the number of visits to x before reaching b is the sum over all y of the number of visits u_y to y before reaching b times the probability p_{yx} of going from yto x. For x not equal to b or a

$$u_x = \sum_{y \neq b} u_y p_{yx}.$$

Since $u_b = 0$ and $c_x p_{xy} = c_y p_{yx}$

$$u_x = \sum_{\text{all } y} u_y \frac{c_x p_{xy}}{c_y}$$

and hence $\frac{u_x}{c_x} = \sum_y \frac{u_y}{c_y} p_{xy}$. It follows that $\frac{u_x}{c_x}$ is harmonic with a and b as the boundary where the boundary conditions are $u_b = 0$ and u_a equals some fixed value. Now, $\frac{u_b}{c_b} = 0$. Setting the current into a to one, fixed the value of v_a . Adjust the current into a so that v_a equals $\frac{u_a}{c_a}$. Now $\frac{u_x}{c_x}$ and v_x satisfy the same boundary conditions and thus are the same harmonic function. Let the current into a correspond to one walk. Note that if the walk starts at a and ends at b, the expected value of the difference between the number of times the walk leaves a and enters a must be one. This implies that the amount of current into a corresponds to one walk.

Next we need to show that the current i_{xy} is the net frequency with which a random walk traverses edge xy.

$$i_{xy} = (v_x - v_y)c_{xy} = \left(\frac{u_x}{c_x} - \frac{u_y}{c_y}\right)c_{xy} = u_x \frac{c_{xy}}{c_x} - u_y \frac{c_{xy}}{c_y} = u_x p_{xy} - u_y p_{yx}$$

The quantity $u_x p_{xy}$ is the expected number of times the edge xy is traversed from x to y and the quantity $u_y p_{yx}$ is the expected number of times the edge xy is traversed from y to x. Thus, the current i_{xy} is the expected net number of traversals of the edge xy from x to y.

Effective resistance and escape probability

Set $v_a = 1$ and $v_b = 0$. Let i_a be the current flowing into the network at vertex a and out at vertex b. Define the *effective resistance* r_{eff} between a and b to be $r_{eff} = \frac{v_a}{i_a}$ and the *effective conductance* c_{eff} to be $c_{eff} = \frac{1}{r_{eff}}$. Define the *escape probability*, p_{escape} , to be the probability that a random walk starting at a reaches b before returning to a. We now show that the escape probability is $\frac{c_{eff}}{c_a}$. For convenience, assume that a and b are not adjacent. A slight modification of the argument suffices for the case when a and b are adjacent.

$$i_a = \sum_y \left(v_a - v_y \right) c_{ay}$$

Since $v_a = 1$,

$$i_a = \sum_y c_{ay} - c_a \sum_y v_y \frac{c_{ay}}{c_a}$$
$$= c_a \left[1 - \sum_y p_{ay} v_y \right].$$

For each y adjacent to the vertex a, p_{ay} is the probability of the walk going from vertex a to vertex y. Earlier we showed that v_y is the probability of a walk starting at y going to a before reaching b. Thus, $\sum_{y} p_{ay}v_y$ is the probability of a walk starting at a returning to a before reaching b and $1 - \sum_{y} p_{ay}v_y$ is the probability of a walk starting at a reaching b before returning to a. Thus, $i_a = c_a p_{escape}$. Since $v_a = 1$ and $c_{eff} = \frac{i_a}{v_a}$, it follows that $c_{eff} = i_a$. Thus, $c_{eff} = c_a p_{escape}$ and hence $p_{escape} = \frac{c_{eff}}{c_a}$.

For a finite connected graph, the escape probability will always be nonzero. Now consider an infinite graph such as a lattice and a random walk starting at some vertex a. Form a series of finite graphs by merging all vertices at distance d or greater from a into a single vertex b for larger and larger values of d. The limit of p_{escape} as d goes to infinity is the probability that the random walk will never return to a. If $p_{escape} \rightarrow 0$, then eventually any random walk will return to a. If $p_{escape} \rightarrow q$ where q > 0, then a fraction of the walks never return. Thus, the escape probability terminology.

5.6 Random Walks on Undirected Graphs with Unit Edge Weights

We now focus our discussion on random walks on undirected graphs with uniform edge weights. At each vertex, the random walk is equally likely to take any edge. This corresponds to an electrical network in which all edge resistances are one. Assume the graph is connected. We consider questions such as what is the expected time for a random walk starting at a vertex x to reach a target vertex y, what is the expected time until the random walk returns to the vertex it started at, and what is the expected time to reach every vertex?

Hitting time

The hitting time h_{xy} , sometimes called discovery time, is the expected time of a random walk starting at vertex x to reach vertex y. Sometimes a more general definition is given where the hitting time is the expected time to reach a vertex y from a given starting probability distribution.

One interesting fact is that adding edges to a graph may either increase or decrease h_{xy} depending on the particular situation. Adding an edge can shorten the distance from x to y thereby decreasing h_{xy} or the edge could increase the probability of a random walk going to some far off portion of the graph thereby increasing h_{xy} . Another interesting fact is that hitting time is not symmetric. The expected time to reach a vertex y from a vertex x in an undirected graph may be radically different from the time to reach x from y.

We start with two technical lemmas. The first lemma states that the expected time to traverse a path of n vertices is $\Theta(n^2)$.

Lemma 5.7 The expected time for a random walk starting at one end of a path of n vertices to reach the other end is $\Theta(n^2)$.

Proof: Consider walking from vertex 1 to vertex n in a graph consisting of a single path of n vertices. Let h_{ij} , i < j, be the hitting time of reaching j starting from i. Now $h_{12} = 1$ and

$$h_{i,i+1} = \frac{1}{2} + \frac{1}{2}(1 + h_{i-1,i+1}) = 1 + \frac{1}{2}(h_{i-1,i} + h_{i,i+1}) \quad 2 \le i \le n-1.$$

Solving for $h_{i,i+1}$ yields the recurrence

$$h_{i,i+1} = 2 + h_{i-1,i}.$$

Solving the recurrence yields

$$h_{i,i+1} = 2i - 1.$$

To get from 1 to n, go from 1 to 2, 2 to 3, etc. Thus

$$h_{1,n} = \sum_{i=1}^{n-1} h_{i,i+1} = \sum_{i=1}^{n-1} (2i-1)$$
$$= 2\sum_{i=1}^{n-1} i - \sum_{i=1}^{n-1} 1$$
$$= 2\frac{n(n-1)}{2} - (n-1)$$
$$= (n-1)^2.$$

The lemma says that in a random walk on a line where we are equally likely to take one step to the right or left each time, the farthest we will go away from the start in nsteps is $\Theta(\sqrt{n})$.

The next lemma shows that the expected time spent at vertex i by a random walk from vertex 1 to vertex n in a chain of n vertices is 2(i-1) for $2 \le i \le n-1$.

Lemma 5.8 Consider a random walk from vertex 1 to vertex n in a chain of n vertices. Let t(i) be the expected time spent at vertex i. Then

$$t(i) = \begin{cases} n-1 & i=1\\ 2(n-i) & 2 \le i \le n-1\\ 1 & i=n. \end{cases}$$

Proof: Now t(n) = 1 since the walk stops when it reaches vertex n. Half of the time when the walk is at vertex n - 1 it goes to vertex n. Thus t(n - 1) = 2. For $3 \le i < n - 1$, $t(i) = \frac{1}{2}[t(i-1) + t(i+1)]$ and t(1) and t(2) satisfy $t(1) = \frac{1}{2}t(2) + 1$ and $t(2) = t(1) + \frac{1}{2}t(3)$. Solving for t(i+1) for $3 \le i < n - 1$ yields

$$t(i+1) = 2t(i) - t(i-1)$$

which has solution t(i) = 2(n-i) for $3 \le i < n-1$. Then solving for t(2) and t(1) yields t(2) = 2(n-2) and t(1) = n-1. Thus, the total time spent at vertices is

$$n-1+2(1+2+\dots+n-2)+1 = (n-1)+2\frac{(n-1)(n-2)}{2}+1 = (n-1)^2+1$$

which is one more than h_{1n} and thus is correct.

Adding edges to a graph might either increase or decrease the hitting time h_{xy} . Consider the graph consisting of a single path of n vertices. Add edges to this graph to get the graph in Figure 5.7 consisting of a clique of size n/2 connected to a path of n/2 vertices. Then add still more edges to get a clique of size n. Let x be the vertex at the midpoint of the original path and let y be the other endpoint of the path consisting of n/2 vertices as shown in the figure. In the first graph consisting of a single path of length n, $h_{xy} = \Theta(n^2)$. In the second graph consisting of a clique of size n/2 along with a path of length n/2, $h_{xy} = \Theta(n^3)$. To see this latter statement, note that starting at x, the walk will go down the path towards y and return to x n/2 times on average before reaching y for the first time. Each time the walk in the path returns to x, with probability (n/2 - 1)/(n/2) it enters the clique and thus on average enters the clique $\Theta(n)$ times before starting down the path again. Each time it enters the clique, it spends $\Theta(n)$ time in the clique before returning to x. Thus, each time the walk returns to x from the path it spends $\Theta(n^2)$ time in the clique before starting down the path towards y for a total expected time that is $\Theta(n^3)$ before reaching y. In the third graph, which is the clique of size n, $h_{xy} = \Theta(n)$. Thus, adding edges first increased h_{xy} from n^2 to n^3 and then decreased it to n.

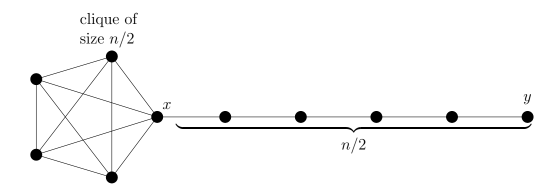


Figure 5.7: Illustration that adding edges to a graph can either increase or decrease hitting time.

Hitting time is not symmetric even in the case of undirected graphs. In the graph of Figure 5.7, the expected time, h_{xy} , of a random walk from x to y, where x is the vertex of attachment and y is the other end vertex of the chain, is $\Theta(n^3)$. However, h_{yx} is $\Theta(n^2)$.

Commute time

The commute time, commute(x, y), is the expected time of a random walk starting at x reaching y and then returning to x. So commute $(x, y) = h_{xy} + h_{yx}$. Think of going from home to office and returning home. We now relate the commute time to an electrical quantity, the effective resistance. The effective resistance between two vertices x and y in an electrical network is the voltage difference between x and y when one unit of current is inserted at vertex x and withdrawn from vertex y.

Theorem 5.9 Given an undirected graph, consider the electrical network where each edge of the graph is replaced by a one ohm resistor. Given vertices x and y, the commute time, commute(x, y), equals $2mr_{xy}$ where r_{xy} is the effective resistance from x to y and m is the number of edges in the graph.

Proof: Insert at each vertex i a current equal to the degree d_i of vertex i. The total current inserted is 2m where m is the number of edges. Extract from a specific vertex j all of this 2m current. Let v_{ij} be the voltage difference from i to j. The current into i divides into the d_i resistors at vertex i. The current in each resistor is proportional to the voltage across it. Let k be a vertex adjacent to i. Then the current through the resistor between i and k is $v_{ij} - v_{kj}$, the voltage drop across the resister. The sum of the currents out of i through the resisters must equal d_i , the current injected into i.

$$d_i = \sum_{\substack{k \text{ adj} \\ \text{to } i}} \left(v_{ij} - v_{kj} \right) = d_i v_{ij} - \sum_{\substack{k \text{ adj} \\ \text{to } i}} v_{kj}.$$

Solving for v_{ij}

$$v_{ij} = 1 + \sum_{\substack{k \text{ adj} \\ \text{to } i}} \frac{1}{d_i} v_{kj} = \sum_{\substack{k \text{ adj} \\ \text{to } i}} \frac{1}{d_i} (1 + v_{kj}).$$
(5.11)

Now the hitting time from i to j is the average time over all paths from i to k adjacent to i and then on from k to j. This is given by

$$h_{ij} = \sum_{\substack{k \text{ adj} \\ \text{to } i}} \frac{1}{d_i} (1 + h_{kj}).$$
(5.12)

Subtracting (5.12) from (5.11), gives $v_{ij} - h_{ij} = \sum_{\substack{k \text{ adj} \\ \text{to } i}} \frac{1}{d_i} (v_{kj} - h_{kj})$. Thus, the function $v_{ij} - h_{ij}$ is harmonic. Designate vertex j as the only boundary vertex. The value of $v_{ij} - h_{ij}$ at i = j, namely $v_{jj} - h_{jj}$, is zero, since both v_{jj} and h_{jj} are zero. So the function $v_{ij} - h_{ij}$ must be zero everywhere. Thus, the voltage v_{ij} equals the expected time h_{ij} from i to j.

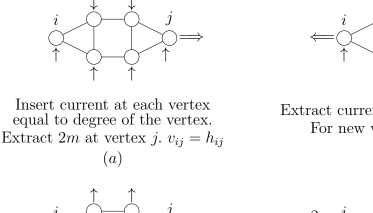
To complete the proof, note that $h_{ij} = v_{ij}$ is the voltage from *i* to *j* when currents are inserted at all vertices in the graph and extracted at vertex *j*. If the current is extracted from *i* instead of *j*, then the voltages change and $v_{ji} = h_{ji}$ in the new setup. Finally, reverse all currents in this latter step. The voltages change again and for the new voltages $-v_{ji} = h_{ji}$. Since $-v_{ji} = v_{ij}$, we get $h_{ji} = v_{ij}$.

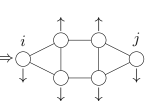
Thus, when a current is inserted at each vertex equal to the degree of the vertex and the current is extracted from j, the voltage v_{ij} in this set up equals h_{ij} . When we extract the current from i instead of j and then reverse all currents, the voltage v_{ij} in this new set up equals h_{ji} . Now, superpose both situations, i.e., add all the currents and voltages. By linearity, for the resulting v_{ij} , which is the sum of the other two v_{ij} 's, is $v_{ij} = h_{ij} + h_{ji}$. All currents cancel except the 2m amps injected at i and withdrawn at j. Thus, $2mr_{ij} = v_{ij} = h_{ij} + h_{ji} = \text{commute}(i, j)$ or $\text{commute}(i, j) = 2mr_{ij}$ where r_{ij} is the effective resistance from i to j.

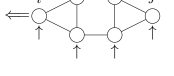
The following corollary follows from Theorem 5.9 since the effective resistance r_{uv} is less than or equal to one when u and v are connected by an edge.

Corollary 5.10 If vertices x and y are connected by an edge, then $h_{xy} + h_{yx} \leq 2m$ where m is the number of edges in the graph.

Proof: If x and y are connected by an edge, then the effective resistance r_{xy} is less than or equal to one.

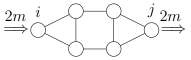






Extract current from *i* instead of *j*. For new voltages $v_{ji} = h_{ji}$.





Reverse currents in (b). For new voltages $-v_{ji} = h_{ji}$. Since $-v_{ji} = v_{ij}, h_{ji} = v_{ij}$. (c)

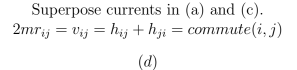


Figure 5.8: Illustration of proof that $commute(x, y) = 2mr_{xy}$ where m is the number of edges in the undirected graph and r_{xy} is the effective resistance between x and y.

Corollary 5.11 For vertices x and y in an n vertex graph, the commute time, commute(x, y), is less than or equal to n^3 .

Proof: By Theorem 5.9 the commute time is given by the formula $\operatorname{commute}(x, y) = 2mr_{xy}$ where *m* is the number of edges. In an *n* vertex graph there exists a path from *x* to *y* of length at most *n*. Since the resistance can not be greater than that of any path from *x* to *y*, $r_{xy} \leq n$. Since the number of edges is at most $\binom{n}{2}$

commute
$$(x, y) = 2mr_{xy} \le 2\binom{n}{2}n \cong n^3.$$

Again adding edges to a graph may increase or decrease the commute time. To see this consider three graphs: the graph consisting of a chain of n vertices, the graph of Figure 5.7, and the clique on n vertices.

Cover time

The cover time, cover(x, G), is the expected time of a random walk starting at vertex x in the graph G to reach each vertex at least once. We write cover(x) when G is understood.

The cover time of an undirected graph G, denoted cover(G), is

$$\operatorname{cover}(G) = \max \operatorname{cover}(x, G)$$

For cover time of an undirected graph, increasing the number of edges in the graph may increase or decrease the cover time depending on the situation. Again consider three graphs, a chain of length n which has cover time $\Theta(n^2)$, the graph in Figure 5.7 which has cover time $\Theta(n^3)$, and the complete graph on n vertices which has cover time $\Theta(n \log n)$. Adding edges to the chain of length n to create the graph in Figure 5.7 increases the cover time from n^2 to n^3 and then adding even more edges to obtain the complete graph reduces the cover time to $n \log n$.

Note: The cover time of a clique is $\theta(n \log n)$ since this is the time to select every integer out of *n* integers with high probability, drawing integers at random. This is called the *coupon collector problem*. The cover time for a straight line is $\Theta(n^2)$ since it is the same as the hitting time. For the graph in Figure 5.7, the cover time is $\Theta(n^3)$ since one takes the maximum over all start states and $\operatorname{cover}(x, G) = \Theta(n^3)$ where x is the vertex of attachment.

Theorem 5.12 Let G be a connected graph with n vertices and m edges. The time for a random walk to cover all vertices of the graph G is bounded above by 4m(n-1).

Proof: Consider a depth first search of the graph G starting from some vertex z and let T be the resulting depth first search spanning tree of G. The depth first search covers every vertex. Consider the expected time to cover every vertex in the order visited by the depth first search. Clearly this bounds the cover time of G starting from vertex z. Note that each edge in T is traversed twice, once in each direction.

$$\operatorname{cover}\left(z,G\right) \le \sum_{\substack{(x,y)\in T\\(y,x)\in T}} h_{xy}.$$

If (x, y) is an edge in T, then x and y are adjacent and thus Corollary 5.10 implies $h_{xy} \leq 2m$. Since there are n-1 edges in the dfs tree and each edge is traversed twice, once in each direction, $\operatorname{cover}(z) \leq 4m(n-1)$. This holds for all starting vertices z. Thus, $\operatorname{cover}(G) \leq 4m(n-1)$.

The theorem gives the correct answer of n^3 for the n/2 clique with the n/2 tail. It gives an upper bound of n^3 for the *n*-clique where the actual cover time is $n \log n$.

Let r_{xy} be the effective resistance from x to y. Define the resistance $r_{eff}(G)$ of a graph G by $r_{eff}(G) = \max_{x,y}(r_{xy})$.

Theorem 5.13 Let G be an undirected graph with m edges. Then the cover time for G is bounded by the following inequality

$$mr_{eff}(G) \le cover(G) \le 2e^3 mr_{eff}(G) \ln n + n$$

where e=2.71 is Euler's constant and $r_{eff}(G)$ is the resistance of G.

Proof: By definition $r_{eff}(G) = \max_{x,y}(r_{xy})$. Let u and v be the vertices of G for which r_{xy} is maximum. Then $r_{eff}(G) = r_{uv}$. By Theorem 5.9, commute $(u, v) = 2mr_{uv}$. Hence $mr_{uv} = \frac{1}{2}$ commute(u, v). Clearly the commute time from u to v and back to u is less than twice the $\max(h_{uv}, h_{vu})$. Finally, $\max(h_{uv}, h_{vu})$ is less than $\max(\operatorname{cover}(u, G), \operatorname{cover}(v, G))$ which is clearly less than the cover time of G. Putting these facts together gives the first inequality in the theorem.

$$mr_{eff}(G) = mr_{uv} = \frac{1}{2} \text{commute}(u, v) \le \max(h_{uv}, h_{vu}) \le \text{cover}(G)$$

For the second inequality in the theorem, by Theorem 5.9, for any x and y, commute(x, y) equals $2mr_{xy}$ which is less than or equal to $2mr_{eff}(G)$, implying $h_{xy} \leq 2mr_{eff}(G)$. By the Markov inequality, since the expected time to reach y starting at any x is less than $2mr_{eff}(G)$, the probability that y is not reached from x in $2mr_{eff}(G)e^3$ steps is at most $\frac{1}{e^3}$. Thus, the probability that a vertex y has not been reached in $2e^3mr_{eff}(G)\log n$ steps is at most $\frac{1}{e^3}\ln^n = \frac{1}{n^3}$ because a random walk of length $2e^3mr(G)\log n$ is a sequence of log n independent random walks, each of length $2e^3mr(G)r_{eff}(G)$. Suppose after a walk of $2e^3mr_{eff}(G)\log n$ steps, vertices v_1, v_2, \ldots, v_l had not been reached. Walk until v_1 is reached, then v_2 , etc. By Corollary 5.11 the expected time for each of these is n^3 , but since each happens only with probability $1/n^3$, we effectively take O(1) time per v_i , for a total time at most n. More precisely,

$$\operatorname{cover}(G) \leq 2e^{3}mr_{eff}(G)\log n + \sum_{v} \operatorname{Prob}\left(v \text{ was not visited in the first } 2e^{3}mr_{eff}(G)\operatorname{steps}\right)n^{3}$$
$$\leq 2e^{3}mr_{eff}(G)\log n + \sum_{v}\frac{1}{n^{3}}n^{3} \leq 2e^{3}mr_{eff}(G) + n.$$

5.7 Random Walks in Euclidean Space

Many physical processes such as Brownian motion are modeled by random walks. Random walks in Euclidean *d*-space consisting of fixed length steps parallel to the coordinate axes are really random walks on a *d*-dimensional lattice and are a special case of random walks on graphs. In a random walk on a graph, at each time unit an edge from the current vertex is selected at random and the walk proceeds to the adjacent vertex. We begin by studying random walks on lattices.

Random walks on lattices

We now apply the analogy between random walks and current to lattices. Consider a random walk on a finite segment $-n, \ldots, -1, 0, 1, 2, \ldots, n$ of a one dimensional lattice starting from the origin. Is the walk certain to return to the origin or is there some probability that it will escape, i.e., reach the boundary before returning? The probability of reaching the boundary before returning to the origin is called the escape probability. We shall be interested in this quantity as n goes to infinity.

Convert the lattice to an electrical network by replacing each edge with a one ohm resister. Then the probability of a walk starting at the origin reaching n or -n before returning to the origin is the escape probability given by

$$p_{escape} = \frac{c_{eff}}{c_a}$$

where c_{eff} is the effective conductance between the origin and the boundary points and c_a is the sum of the conductance's at the origin. In a *d*-dimensional lattice, $c_a = 2d$ assuming that the resistors have value one. For the *d*-dimensional lattice

$$p_{escape} = \frac{1}{2d \ r_{eff}}$$

In one dimension, the electrical network is just two series connections of n one ohm resistors connected in parallel. So as n goes to infinity, r_{eff} goes to infinity and the escape probability goes to zero as n goes to infinity. Thus, the walk in the unbounded one dimensional lattice will return to the origin with probability one. This is equivalent to flipping a balanced coin and keeping tract of the number of heads minus the number of tails. The count will return to zero infinitely often.

Two dimensions

For the 2-dimensional lattice, consider a larger and larger square about the origin for the boundary as shown in Figure 5.9a and consider the limit of r_{eff} as the squares get larger. Shorting the resistors on each square can only reduce r_{eff} . Shorting the resistors results in the linear network shown in Figure 5.9b. As the paths get longer, the number of resistors in parallel also increases. The resistor between vertex i and i + 1 is really 4(2i+1) unit resistors in parallel. The effective resistance of 4(2i+1) resistors in parallel is 1/4(2i+1). Thus,

$$r_{eff} \ge \frac{1}{4} + \frac{1}{12} + \frac{1}{20} + \dots = \frac{1}{4}(1 + \frac{1}{3} + \frac{1}{5} + \dots) = \Theta(\ln n).$$

Since the lower bound on the effective resistance and hence the effective resistance goes to infinity, the escape probability goes to zero for the 2-dimensional lattice.

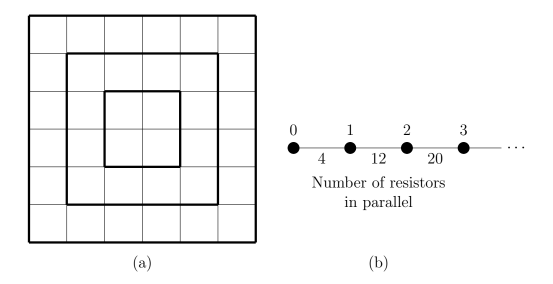


Figure 5.9: 2-dimensional lattice along with the linear network resulting from shorting resistors on the concentric squares about the origin.

Three dimensions

In three dimensions, the resistance along any path to infinity grows to infinity but the number of paths in parallel also grows to infinity. It turns out there are a sufficient number of paths that r_{eff} remains finite and thus there is a nonzero escape probability. We will prove this now. First note that shorting any edge decreases the resistance, so we do not use shorting in this proof, since we seek to prove an upper bound on the resistance. Instead we remove some edges, which increases their resistance to infinity and hence increases the effective resistance, giving an upper bound. To simplify things we consider walks on on quadrant rather than the full grid. The resistance to infinity derived from only the quadrant is an upper bound on the resistance of the full grid.

The construction used in three dimensions is easier to explain first in two dimensions. Draw dotted diagonal lines at $x + y = 2^n - 1$. Consider two paths that start at the origin. One goes up and the other goes to the right. Each time a path encounters a dotted diagonal line, split the path into two, one which goes right and the other up. Where two paths cross, split the vertex into two, keeping the paths separate. By a symmetry argument, splitting the vertex does not change the resistance of the network. Remove all resistors except those on these paths. The resistance of the original network is less than that of the tree produced by this process since removing a resistor is equivalent to increasing its resistance to infinity.

The distances between splits increase and are 1, 2, 4, etc. At each split the number of paths in parallel doubles. See Figure 5.11. Thus, the resistance to infinity in this two

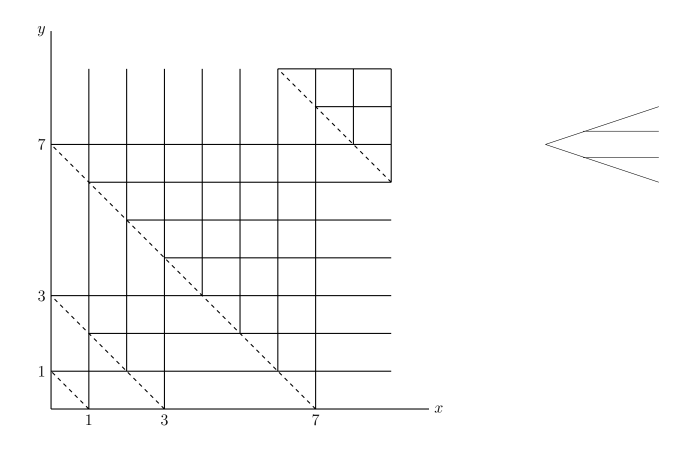


Figure 5.10: Paths in a 2-dimensional lattice obtained from the 3-dimensional construction applied in 2-dimensions.

dimensional example is

$$\frac{1}{2} + \frac{1}{4}2 + \frac{1}{8}4 + \dots = \frac{1}{2} + \frac{1}{2} + \frac{1}{2} + \dots = \infty.$$

In the analogous three dimensional construction, paths go up, to the right, and out of the plane of the paper. The paths split three ways at planes given by $x + y + z = 2^n - 1$. Each time the paths split the number of parallel segments triple. Segments of the paths between splits are of length 1, 2, 4, etc. and the resistance of the segments are equal to the lengths. The resistance out to infinity for the tree is

$$\frac{1}{3} + \frac{1}{9}2 + \frac{1}{27}4 + \dots = \frac{1}{3}\left(1 + \frac{2}{3} + \frac{4}{9} + \dots\right) = \frac{1}{3}\frac{1}{1 - \frac{2}{3}} = 1$$

The resistance of the three dimensional lattice is less. It is important to check that the paths are edge-disjoint and so the tree is a subgraph of the lattice. Going to a subgraph is equivalent to deleting edges which only increases the resistance. That is why the resistance

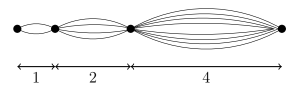


Figure 5.11: Paths obtained from 2-dimensional lattice. Distances between splits double as do the number of parallel paths.

of the lattice is less than that of the tree. Thus, in three dimensions the escape probability is nonzero. The upper bound on r_{eff} gives the lower bound

$$p_{escape} = \frac{1}{2d} \frac{1}{r_{eff}} \ge \frac{1}{6}.$$

A lower bound on r_{eff} gives an upper bound on p_{escape} . To get the upper bound on p_{escape} , short all resistors on surfaces of boxes at distances 1, 2, 3, etc. Then

$$r_{eff} \ge \frac{1}{6} \left[1 + \frac{1}{9} + \frac{1}{25} + \cdots \right] \ge \frac{1.23}{6} \ge 0.2$$

This gives

$$p_{escape} = \frac{1}{2d} \frac{1}{r_{eff}} \le \frac{5}{6}.$$

5.8 The Web as a Markov Chain

A modern application of random walks on directed graphs comes from trying to establish the importance of pages on the World Wide Web. Search Engines output an ordered list of webpages in response to each search query. To do this, they have to solve two problems at query time: (i) find the set of all webpages containing the query term(s) and (ii) rank the webpages and display them (or the top subset of them) in ranked order. (i) is done by maintaining a "reverse index" which we do not discuss here. (ii) cannot be done at query time since this would make the response too slow. So Search Engines rank the entire set of webpages (in the billions) "off-line" and use that single ranking for all queries. At query time, the webpages containing the query terms(s) are displayed in this ranked order.

One way to do this ranking would be to take a random walk on the web viewed as a directed graph (which we call the web graph) with an edge corresponding to each hypertext link and rank pages according to their stationary probability. Hypertext links are one-way and the web graph may not be strongly connected. Indeed, for a node at the "bottom" level there may be no out-edges. When the walk encounters this vertex the walk disappears. Another difficulty is that a vertex or a strongly connected component with no in edges is never reached. One way to resolve these difficulties is to introduce a random restart condition. At each step, with some probability r, jump to a vertex selected uniformly at random and with probability 1 - r select an edge at random and follow it. If a vertex has no out edges, the value of r for that vertex is set to one. This makes the

Figure 5.12: Impact on page rank of adding a self loop

graph strongly connected so that the stationary probabilities exist.

Page rank

The page rank of a vertex in a directed graph is the stationary probability of the vertex, where we assume a positive restart probability of say r = 0.15. The restart ensures that the graph is strongly connected. The page rank of a page is the fractional frequency with which the page will be visited over a long period of time. If the page rank is p, then the expected time between visits or return time is 1/p. Notice that one can increase the pagerank of a page by reducing the return time and this can be done by creating short cycles.

Consider a vertex *i* with a single edge in from vertex *j* and a single edge out. The stationary probability $\boldsymbol{\pi}$ satisfies $\boldsymbol{\pi}P = \boldsymbol{\pi}$, and thus

$$\pi_i = \pi_j p_{ji}.$$

Adding a self-loop at i, results in a new equation

$$\pi_i = \pi_j p_{ji} + \frac{1}{2}\pi_i$$

or

$$\pi_i = 2 \ \pi_j p_{ji}.$$

Of course, π_j would have changed too, but ignoring this for now, pagerank is doubled by the addition of a self-loop. Adding k self loops, results in the equation

$$\pi_i = \pi_j p_{ji} + \frac{k}{k+1} \pi_i,$$

and again ignoring the change in π_j , we now have $\pi_i = (k+1)\pi_j p_{ji}$. What prevents one from increasing the page rank of a page arbitrarily? The answer is the restart. We neglected the 0.15 probability that is taken off for the random restart. With the restart taken into account, the equation for π_i when there is no self-loop is

$$\pi_i = 0.85\pi_j p_{ji}$$

whereas, with k self-loops, the equation is

$$\pi_i = 0.85\pi_j p_{ji} + 0.85\frac{k}{k+1}\pi_i.$$

Solving for π_i yields

$$\pi_i = \frac{0.85k + 0.85}{0.15k + 1} \pi_j p_{ji}$$

which for k = 1 is $\pi_i = 1.48\pi_j P_{ji}$ and in the limit as $k \to \infty$ is $\pi_i = 5.67\pi_j p_{ji}$. Adding a single loop only increases pagerank by a factor of 1.74.

Relation to Hitting time

Recall the definition of hitting time h_{xy} , which for two states x, y is the expected time to first reach y starting from x. Here, we deal with H_y , the average time to hit y, starting at a random node. Namely, $H_y = \frac{1}{n} \sum_x h_{xy}$, where the sum is taken over all nodes x. [The number of nodes is n.] Hitting time H_y is closely related to return time and thus to the reciprocal of page rank. Return time is clearly less than the expected time until a restart plus hitting time. This gives:

Return time to
$$y \le .15 \times 1 + .85 \times .15 \times 2 + (.85)^2 \times .15 \times 3 + \dots + H_y \le \frac{1}{.15} + H_y.$$

In the other direction, the fastest one could return would be if there were only paths of length two since self loops are ignored in calculating page rank. If r is the restart value, then the loop would be traversed with at most probability $(1-r)^2$. With probability r + (1-r)r = (2-r)r one restarts and then hits v. Thus, the return time is at least $2(1-r)^2 + (2-r)r \times$ (hitting time). Combining these two bounds yields

$$2(1-r)^2 + (2-r)r(\text{hitting time}) \le (\text{return time}) \le 6.66 + (\text{hitting time}).$$

The relationship between return time and hitting time can be used to see if a vertex has unusually high probability of short loops. However, there is no efficient way to compute hitting time for all vertices as there is for return time. For a single vertex v, one can compute hitting time by removing the edges out of the vertex v for which one is computing hitting time and then run the page rank algorithm for the new graph. The hitting time for v is the reciprocal of the page rank in the graph with the edges out of v removed. Since computing hitting time for each vertex requires removal of a different set of edges, the algorithm only gives the hitting time for one vertex at a time. Since one is probably only interested in the hitting time of vertices with low hitting time, an alternative would be to use a random walk to estimate the hitting time of low hitting time vertices.

Spam

Suppose one has a web page and would like to increase its page rank by creating some other web pages with pointers to the original page. The abstract problem is the following. We are given a directed graph G and a vertex v whose page rank we want to increase. We may add new vertices to the graph and add edges from v or from the new vertices to any vertices we want. We cannot add edges out of other vertices. We can also delete edges from v.

The page rank of v is the stationary probability for vertex v with random restarts. If we delete all existing edges out of v, create a new vertex u and edges (v, u) and (u, v), then the page rank will be increased since any time the random walk reaches v it will be captured in the loop $v \to u \to v$. A search engine can counter this strategy by more frequent random restarts.

A second method to increase page rank would be to create a star consisting of the vertex v at its center along with a large set of new vertices each with a directed edge to v. These new vertices will sometimes be chosen as the target of the random restart and hence the vertices increase the probability of the random walk reaching v. This second method is countered by reducing the frequency of random restarts.

Notice that the first technique of capturing the random walk increases page rank but does not effect hitting time. One can negate the impact of someone capturing the random walk on page rank by increasing the frequency of random restarts. The second technique of creating a star increases page rank due to random restarts and decreases hitting time. One can check if the page rank is high and hitting time is low in which case the page rank is likely to have been artificially inflated by the page capturing the walk with short cycles.

Personalized page rank

In computing page rank, one uses a restart probability, typically 0.15, in which at each step, instead of taking a step in the graph, the walk goes to a vertex selected uniformly at random. In personalized page rank, instead of selecting a vertex uniformly at random, one selects a vertex according to a personalized probability distribution. Often the distribution has probability one for a single vertex and whenever the walk restarts it restarts at that vertex.

Algorithm for computing personalized page rank

First, consider the normal page rank. Let α be the restart probability with which the random walk jumps to an arbitrary vertex. With probability $1 - \alpha$ the random walk selects a vertex uniformly at random from the set of adjacent vertices. Let **p** be a row vector denoting the page rank and let A be the adjacency matrix with rows normalized to sum to one. Then

$$\mathbf{p} = \frac{\alpha}{n} \left(1, 1, \dots, 1 \right) + \left(1 - \alpha \right) \mathbf{p} A$$

$$\mathbf{p}[I - (1 - \alpha)A] = \frac{\alpha}{n}(1, 1, \dots, 1)$$
$$\mathbf{p} = \frac{\alpha}{n}(1, 1, \dots, 1)[I - (1 - \alpha)A]^{-1}.$$

or

$$\mathbf{p} = \frac{\alpha}{n} (1, 1, \dots, 1) \left[I - (1 - \alpha) A \right]^{-1}.$$

Thus, in principle, **p** can be found by computing the inverse of $[I - (1 - \alpha)A]^{-1}$. But this is far from practical since for the whole web one would be dealing with matrices with billions of rows and columns. A more practical procedure is to run the random walk and observe using the basics of the power method in Chapter 3 that the process converges to the solution **p**.

For the personalized page rank, instead of restarting at an arbitrary vertex, the walk restarts at a designated vertex. More generally, it may restart in some specified neighborhood. Suppose the restart selects a vertex using the probability distribution s. Then, in the above calculation replace the vector $\frac{1}{n}(1,1,\ldots,1)$ by the vector s. Again, the computation could be done by a random walk. But, we wish to do the random walk calculation for personalized pagerank quickly since it is to be performed repeatedly. With more care this can be done, though we do not describe it here.

5.9 **Bibliographic Notes**

The material on the analogy between random walks on undirected graphs and electrical networks is from [DS84] as is the material on random walks in Euclidean space. Additional material on Markov chains can be found in [MR95b], [MU05], and [per10]. For material on Markov Chain Monte Carlo methods see [Jer98] and [Liu01].

The use of normalized conductance to prove convergence of Markov Chains is by Sinclair and Jerrum, [SJ] and Alon [Alo86]. A polynomial time bounded Markov chain based method for estimating the volume of convex sets was developed by Dyer, Frieze and Kannan [DFK91].

5.10 Exercises

Exercise 5.1 The Fundamental Theorem of Markov chains proves that for a connected Markov chain, the long-term average distribution $\mathbf{a}_{\mathbf{t}}$ converges to a stationary distribution. Does the t step distribution $\mathbf{p}_{\mathbf{t}}$ also converge for every connected Markov Chain? Consider the following examples: (i) A two-state chain with $p_{12} = p_{21} = 1$. (ii) A three state chain with $p_{12} = p_{23} = p_{31} = 1$ and the other $p_{ij} = 0$. Generalize these examples to produce Markov Chains with many states.

Exercise 5.2 Let $p(\mathbf{x})$, where $\mathbf{x} = (x_1, x_2, \dots, x_d)$ $x_i \in \{0, 1\}$, be a multivariate probability distribution. For d = 100, how would you estimate the marginal distribution

$$p(x_1) = \sum_{x_2,\dots,x_d} p(x_1, x_2, \dots, x_d)?$$

Exercise 5.3 Prove $|\mathbf{p} - \mathbf{q}|_1 = 2\sum_i (p_i - q_i)^+$ for probability distributions \mathbf{p} and \mathbf{q} . Proposition 5.4

Exercise 5.4 Suppose S is a subset of at most $n^2/2$ points in the $n \times n$ lattice. Show that for

 $T = \{(i, j) \in S | all elements in row i and all elements in column j are in S \}$

 $|T| \le |S|/2.$

Exercise 5.5 Show that the stationary probabilities of the chain described in the Gibbs sampler is the correct p.

Exercise 5.6 A Markov chain is said to be symmetric if for all i and j, $p_{ij} = p_{ji}$. What is the stationary distribution of a connected symmetric chain? Prove your answer.

Exercise 5.7 How would you integrate a high dimensional multivariate polynomial distribution over some convex region?

this exercise needs to be clarified, ARE WE IN HIGH DIMENSIONS, IS REGION CONVEX?

Exercise 5.8 Given a time-reversible Markov chain, modify the chain as follows. At the current state, stay put (no move) with probability 1/2. With the other probability 1/2, move as in the old chain. Show that the new chain has the same stationary distribution. What happens to the convergence time in this modification?

Exercise 5.9 Using the Metropolis-Hasting Algorithm create a Markov chain whose stationary probability is that given in the following table.

x_1x_2	00	01	02	10	11	12	20	21	22
Prob	1/16	1/8	1/16	1/8	1/4	1/8	1/16	1/8	1/16

Exercise 5.10 Let \mathbf{p} be a probability vector (nonnegative components adding up to 1) on the vertices of a connected graph which is sufficiently large that it cannot be stored in a computer. Set p_{ij} (the transition probability from i to j) to p_j for all $i \neq j$ which are adjacent in the graph. Show that the stationary probability vector is \mathbf{p} . Is a random walk an efficient way to sample according to a distribution close to \mathbf{p} ? Think, for example, of the graph G being the $n \times n \times n \times \cdots n$ grid.

Exercise 5.11 Construct the edge probability for a three state Markov chain where each pair of states is connected by an edge so that the stationary probability is $(\frac{1}{2}, \frac{1}{3}, \frac{1}{6})$.

Exercise 5.12 Consider a three state Markov chain with stationary probability $(\frac{1}{2}, \frac{1}{3}, \frac{1}{6})$. Consider the Metropolis-Hastings algorithm with G the complete graph on these three vertices. What is the expected probability that we would actually make a move along a selected edge?

Exercise 5.13 Try Gibbs sampling on $p(x) = \begin{pmatrix} \frac{1}{2} & 0\\ 0 & \frac{1}{2} \end{pmatrix}$. What happens? How does the Metropolis Hasting Algorithm do?

Exercise 5.14 Consider $p(\mathbf{x})$, where, $\mathbf{x} = (x_1, \dots, x_{100})$ and $p(\mathbf{0}) = \frac{1}{2}$, $p(\mathbf{x}) = \frac{1}{(2^{100}-1)}$ $\mathbf{x} \neq \mathbf{0}$. How does Gibbs sampling behave?

Exercise 5.15 Construct, program, and execute an algorithm to compute the volume of a unit radius sphere in 20 dimensions by carrying out a random walk on a 20 dimensional grid with 0.1 spacing.

Exercise 5.16 Given a connected graph G and an integer k how would you generate connected subgraphs of G with k vertices with probability proportional to the number of edges in the subgraph induced on those vertices? The probabilities need not be exactly proportional to the number of edges and you are not expected to prove your algorithm for this problem.

Exercise 5.17 Suppose one wishes to generate uniformly at random regular, degree three undirected, connected multi-graphs each with 1,000 vertices. A multi-graph may have multiple edges between a pair of vertices and self loops. One decides to do this by a Markov Chain Monte Carlo technique. They design a network where each vertex is a regular degree three, 1,000 vertex multi-graph. For edges they say that the vertices corresponding to two graphs are connected by an edge if one graph can be obtained from the other by a flip of a pair of disjoint edges. In a flip, a pair of edges (a, b) and (c, d) are replaced by (a, c) and (b, d).

- 1. Prove that a swap on a connected multi-graph results in a connected multi-graph.
- 2. Prove that the network whose vertices correspond to the desired graphs is connected.

- 3. Prove that the stationary probability of the random walk is uniform. Is there a better word than unifrom?
- 4. Give an upper bound on the diameter of the network.

In order to use a random walk to generate the graphs uniformly at random, the random walk must rapidly converge to the stationary probability. Proving this is beyond the material in this book.

Exercise 5.18 What is the mixing time for

- 1. Two cliques connected by a single edge?
- 2. A graph consisting of an n vertex clique plus one additional vertex connected to one vertex in the clique.

Exercise 5.19 What is the mixing time for

- 1. G(n,p) with $p = \frac{\log n}{n}$?
- 2. A circle with n vertices where at each vertex an edge has been added to another vertex chosen at random. On average each vertex will have degree four, two circle edges, and an edge from that vertex to a vertex chosen at random, and possible some edges that are the ends of the random edges from other vertices.

Exercise 5.20 Show that for the $n \times n \times \cdots \times n$ grid in d space, the normalized conductance is $\Omega(1/dn)$. Hint: The argument is a generalization of the argument in Exercise 5.4. Argue that for any subset S containing at most 1/2 the grid points, for at least 1/2 the grid points in S, among the d coordinate lines through the point, at least one intersects \overline{S} .

Exercise 5.21

- 1. What is the set of possible harmonic functions on a connected graph if there are only interior vertices and no boundary vertices that supply the boundary condition?
- 2. Let q_x be the stationary probability of vertex x in a random walk on an undirected graph where all edges at a vertex are equally likely and let d_x be the degree of vertex x. Show that $\frac{q_x}{d_x}$ is a harmonic function.
- 3. If there are multiple harmonic functions when there are no boundary conditions, why is the stationary probability of a random walk on an undirected graph unique?
- 4. What is the stationary probability of a random walk on an undirected graph?

Exercise 5.22 In Section ?? we associate a graph and edge probabilities with an electric network such that voltages and currents in the electrical network corresponded to properties of random walks on the graph. Can we go in the reverse order and construct the equivalent electrical network from a graph with edge probabilities?

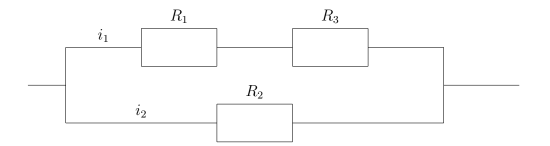


Figure 5.13: An electrical network of resistors.

Exercise 5.23 Given an undirected graph consisting of a single path of five vertices numbered 1 to 5, what is the probability of reaching vertex 1 before vertex 5 when starting at vertex 4.

Exercise 5.24 Consider the electrical resistive network in Figure 5.13 consisting of vertices connected by resistors. Kirchoff's law states that the currents at each vertex sum to zero. Ohm's law states that the voltage across a resistor equals the product of the resistance times the current through it. Using these laws calculate the effective resistance of the network.

Exercise 5.25 Consider the electrical network of Figure 5.14.

- 1. Set the voltage at a to one and at b to zero. What are the voltages at c and d?
- 2. What is the current in the edges a to c, a to d, c to d. c to b and d to b?
- 3. What is the effective resistance between a and b?
- 4. Convert the electrical network to a graph. What are the edge probabilities at each vertex?
- 5. What is the probability of a walk starting at c reaching a before b? a walk starting at d reaching a before b?
- 6. What is the net frequency that a walk from a to b goes through the edge from c to d?
- 7. What is the probability that a random walk starting at a will return to a before reaching b?

Exercise 5.26 Consider a graph corresponding to an electrical network with vertices a and b. Prove directly that $\frac{c_{eff}}{c_a}$ must be less than or equal to one. We know that this is the escape probability and must be at most 1. But, for this exercise, do not use that fact.

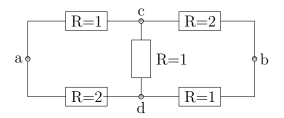


Figure 5.14: An electrical network of resistors.

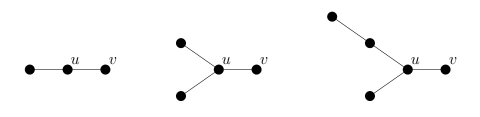


Figure 5.15: Three graphs

Exercise 5.27 (Thomson's Principle) The energy dissipated by the resistance of edge xy in an electrical network is given by $i_{xy}^2 r_{xy}$. The total energy dissipation in the network is $E = \frac{1}{2} \sum_{x,y} i_{xy}^2 r_{xy}$ where the $\frac{1}{2}$ accounts for the fact that the dissipation in each edge is counted twice in the summation. Show that the actual current distribution is the distribution satisfying Ohm's law that minimizes energy dissipation.

Exercise 5.28 (Rayleigh's law) Prove that reducing the value of a resistor in a network cannot increase the effective resistance. Prove that increasing the value of a resistor cannot decrease the effective resistance. You may use Thomson's principle Exercise 5.27.

Exercise 5.29 What is the hitting time h_{uv} for two adjacent vertices on a cycle of length n? What is the hitting time if the edge (u, v) is removed?

Exercise 5.30 What is the hitting time h_{uv} for the three graphs if Figure 5.15.

Exercise 5.31 Show that adding an edge can either increase or decrease hitting time by calculating h_{24} for the three graphs in Figure 5.16.

Exercise 5.32 Consider the n vertex connected graph shown in Figure 5.17 consisting of an edge (u, v) plus a connected graph on n - 1 vertices and m edges. Prove that $h_{uv} = 2m + 1$ where m is the number of edges in the n - 1 vertex subgraph.

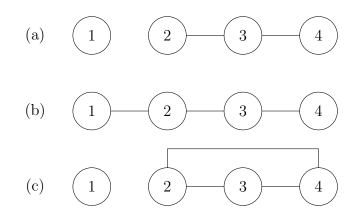


Figure 5.16: Three graph

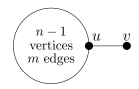


Figure 5.17: A connected graph consisting of n-1 vertices and m edges along with a single edge (u, v).

Exercise 5.33 What is the most general solution to the difference equation t(i + 2) - 5t(i + 1) + 6t(i) = 0. How many boundary conditions do you need to make the solution unique?

Exercise 5.34 Given the difference equation $a_k t(i+k) + a_{k-1}t(i+k-1) + \cdots + a_1t(i+1) + a_0t(i) = 0$ the polynomial $a_k t^k + a_{k-i}t^{k-1} + \cdots + a_1t + a_0 = 0$ is called the characteristic polynomial.

- 1. If the equation has a set of r distinct roots, what is the most general form of the solution?
- 2. If the roots of the characteristic polynomial are not distinct what is the most general form of the solution?
- 3. What is the dimension of the solution space?
- 4. If the difference equation is not homogeneous (i.e., the right hand side is not 0) and f(i) is a specific solution to the nonhomogeneous difference equation, what is the full set of solutions to the difference equation?

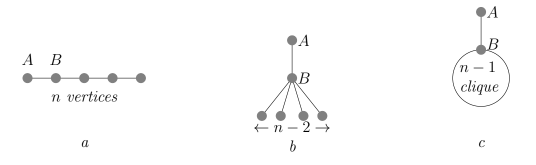
Exercise 5.35 Given the integers 1 to n, what is the expected number of draws with replacement until the integer 1 is drawn.

Exercise 5.36 Consider the set of integers $\{1, 2, ..., n\}$. What is the expected number of draws d with replacement so that every integer is drawn?

Exercise 5.37 Consider a random walk on a clique of size n. What is the expected number of steps before a given vertex is reached?

Exercise 5.38 Show that adding an edge to a graph can either increase or decrease commute time.

Exercise 5.39 For each of the three graphs below what is the return time starting at vertex A? Express your answer as a function of the number of vertices, n, and then express it as a function of the number of edges m.



Exercise 5.40 Suppose that the clique in Exercise 5.39 was replaced by an arbitrary graph with m-1 edges. What would be the return time to A in terms of m, the total number of edges.

Exercise 5.41 Suppose that the clique in Exercise 5.39 was replaced by an arbitrary graph with m-d edges and there were d edges from A to the graph. What would be the expected length of a random path starting at A and ending at A after returning to A exactly d times.

Exercise 5.42 Given an undirected graph with a component consisting of a single edge find two eigenvalues of the Laplacian L = D - A where D is a diagonal matrix with vertex degrees on the diagonal and A is the adjacency matrix of the graph.

Exercise 5.43 A researcher was interested in determining the importance of various edges in an undirected graph. He computed the stationary probability for a random walk on the graph and let p_i be the probability of being at vertex *i*. If vertex *i* was of degree d_i , the frequency that edge (i, j) was traversed from *i* to *j* would be $\frac{1}{d_i}p_i$ and the frequency that the edge was traversed in the opposite direction would be $\frac{1}{d_j}p_j$. Thus, he assigned an importance of $\left|\frac{1}{d_i}p_i - \frac{1}{d_j}p_j\right|$ to the edge. What is wrong with his idea?

Exercise 5.44 Prove that two independent random walks starting at the origin on a two dimensional lattice will eventually meet with probability one.

Exercise 5.45 Suppose two individuals are flipping balanced coins and each is keeping tract of the number of heads minus the number of tails. Will both individual's counts ever return to zero at the same time?

Exercise 5.46 Consider the lattice in 2-dimensions. In each square add the two diagonal edges. What is the escape probability for the resulting graph?

Exercise 5.47 Determine by simulation the escape probability for the 3-dimensional lattice.

Exercise 5.48 What is the escape probability for a random walk starting at the root of an infinite binary tree?

Exercise 5.49 Consider a random walk on the positive half line, that is the integers $0, 1, 2, \ldots$ At the origin, always move right one step. At all other integers move right with probability 2/3 and left with probability 1/3. What is the escape probability?

Exercise 5.50 Consider the graphs in Figure 5.18. Calculate the stationary distribution for a random walk on each graph and the flow through each edge. What condition holds on the flow through edges in the undirected graph? In the directed graph?

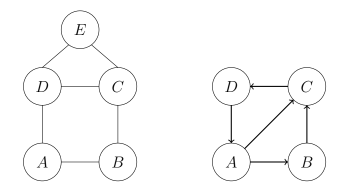


Figure 5.18: An undirected and a directed graph.

Exercise 5.51 Create a random directed graph with 200 vertices and roughly eight edges per vertex. Add k new vertices and calculate the page rank with and without directed edges from the k added vertices to vertex 1. How much does adding the k edges change the page rank of vertices for various values of k and restart frequency? How much does adding a loop at vertex 1 change the page rank? To do the experiment carefully one needs to consider the page rank of a vertex to which the star is attached. If it has low page rank its page rank is likely to increase a lot.

Exercise 5.52 Repeat the experiment in Exercise 5.51 for hitting time.

Exercise 5.53 Search engines ignore self loops in calculating page rank. Thus, to increase page rank one needs to resort to loops of length two. By how much can you increase the page rank of a page by adding a number of loops of length two?

Exercise 5.54 Number the vertices of a graph $\{1, 2, ..., n\}$. Define hitting time to be the expected time from vertex 1. In (2) assume that the vertices in the cycle are sequentially numbered.

- 1. What is the hitting time for a vertex in a complete directed graph with self loops?
- 2. What is the hitting time for a vertex in a directed cycle with n vertices?

Create exercise relating strongly connected and full rank Full rank implies strongly connected. Strongly connected does not necessarily imply full rank

$$\left(\begin{array}{rrr} 0 & 0 & 1 \\ 0 & 0 & 1 \\ 1 & 1 & 0 \end{array}\right)$$

Is graph aperiodic iff $\lambda_1 > \lambda_2$?

Exercise 5.55 Using a web browser bring up a web page and look at the source html. How would you extract the url's of all hyperlinks on the page if you were doing a crawl of the web? With Internet Explorer click on "source" under "view" to access the html representation of the web page. With Firefox click on "page source" under "view".

Exercise 5.56 Sketch an algorithm to crawl the World Wide Web. There is a time delay between the time you seek a page and the time you get it. Thus, you cannot wait until the page arrives before starting another fetch. There are conventions that must be obeyed if one were to actually do a search. Sites specify information as to how long or which files can be searched. Do not attempt an actual search without guidance from a knowledgeable person.

6 Machine Learning

6.1 Introduction

Machine learning algorithms are general purpose tools that solve problems from many disciplines without detailed domain-specific knowledge. They have proven to be very effective in a large number of contexts, including computer vision, speech recognition, document classification, automated driving, computational science, and decision support.

The core problem. A core problem underlying many machine learning applications is learning a good classification rule from labeled data. This problem consists of a domain of interest \mathcal{X} , called the *instance space*, such as email messages or patient records, and a classification task, such as classifying email messages into spam versus non-spam or determining which patients will respond well to a given medical treatment. We will typically assume our instance space $\mathcal{X} = \{0, 1\}^d$ or $\mathcal{X} = \mathbb{R}^d$, corresponding to data that is described by d Boolean or real-valued features. Features for email messages could be the presence or absence of various types of words, and features for patient records could be the results of various medical tests. To perform the learning task, our learning algorithm is given a set S of labeled *training examples*, which are points in \mathcal{X} along with their correct classification. This training data could be a collection of email messages, each labeled as spam or not spam, or a collection of patients, each labeled by whether or not they responded well to the given medical treatment. Our algorithm then aims to use the training examples to produce a classification rule that will perform well over new data. A key feature of machine learning, which distinguishes it from other algorithmic tasks, is that our goal is generalization: to use one set of data in order to perform well on new data we have not seen yet. We focus on *binary classification* where items in the domain of interest are classified into two categories, as in the medical and spam-detection examples above.

How to learn. A high-level approach to solving this problem that many algorithms we discuss will follow is to try to find a "simple" rule with good performance on the training data. For instance in the case of classifying email messages, we might find a set of highly indicative words such that every spam email in the training data has at least one of these words and none of the non-spam emails has any of them; in this case, the rule "if the message has any of these words then it is spam, else it is not" would be a simple rule that performs well on the training data. Or, we might find a way of weighting words with positive and negative weights such that the total weighted sum of words in the email message is positive on the spam emails in the training data, and negative on the non-spam emails. We will then argue that so long as the training data is representative of what future data will look like, we can be confident that any sufficiently "simple" rule that performs well on the training data to be precise about what we mean by "simple" as well as what it means for training data to be "representative" of future data. In fact, we will see several notions of complexity, including bit-counting and VC-dimension, that

will allow us to make mathematical statements of this form. These statements can be viewed as formalizing the intuitive philosophical notion of Occam's razor.

Formalizing the problem. To formalize the learning problem, assume there is some probability distribution D over the instance space \mathcal{X} , such that (a) our training set S consists of points drawn independently at random from D, and (b) our objective is to predict well on new points that are also drawn from D. This is the sense in which we assume that our training data is representative of future data. Let c^* , called the *target* concept, denote the subset of \mathcal{X} corresponding to the positive class for the binary classification we are aiming to make. For example, c^* would correspond to the set of all patients who respond well to the treatment in the medical example, or the set of all spam emails in the spam-detection setting. So, each point in our training set S is labeled according to whether or not it belongs to c^* and our goal is to produce a set $h \subset \mathcal{X}$, called our hypothesis, which is close to c^* with respect to distribution D. The true error of h is $err_D(h) = \operatorname{Prob}(h \triangle c^*)$ where " \triangle " denotes symmetric difference, and probability mass is according to D. In other words, the true error of h is the probability it incorrectly classifies a data point drawn at random from D. Our goal is to produce h of low true error. The training error of h, denoted $err_S(h)$, is the fraction of points in S on which h and c^* disagree. That is, $err_S(h) = |S \cap (h \triangle c^*)| / |S|$. Training error is also called *empirical* error. Note that even though S is assumed to consist of points randomly drawn from D, it is possible for a hypothesis h to have low training error or even to completely agree with c^* over the training sample, and yet have high true error. This is called *overfitting* the training data. For instance, a hypothesis h that simply consists of listing the positive examples in S, which is equivalent to a rule that memorizes the training sample and predicts positive on an example if and only if it already appeared positively in the training sample, would have zero training error. However, this hypothesis likely would have high true error and therefore would be highly overfitting the training data. More generally, overfitting is a concern because algorithms will typically be optimizing over the training sample. To design and analyze algorithms for learning, we will have to address the issue of overfitting.

To be able to formally analyze overfitting, we introduce the notion of an hypothesis class, also called a concept class or set system. An hypothesis class \mathcal{H} over \mathcal{X} is a collection of subsets of \mathcal{X} , called hypotheses. For instance, the class of *intervals* over $\mathcal{X} = \mathbb{R}$ is the collection $\{[a, b] | a \leq b\}$. The class of *linear separators* over $\mathcal{X} = \mathbb{R}^d$ is the collection

$$\{\{\mathbf{x} \in \mathbb{R}^d | \mathbf{w} \cdot \mathbf{x} \ge w_0\} | \mathbf{w} \in \mathbb{R}^d, w_0 \in \mathbb{R}\};\$$

that is, it is the collection of all sets in \mathbb{R}^d that are linearly separable from their complement. In the case that \mathcal{X} is the set of 4 points in the plane $\{(-1, -1), (-1, 1), (1, -1), (1, 1)\}$, the class of linear separators contains 14 of the $2^4 = 16$ possible subsets of \mathcal{X} .¹⁷ Given an hypothesis class \mathcal{H} and training set S, what we typically aim to do algorithmically is to find the hypothesis in \mathcal{H} that most closely agrees with c^* over S. To address overfitting,

¹⁷The only two subsets that are not in the class are the sets $\{(-1, -1), (1, 1)\}$ and $\{(-1, 1), (1, -1)\}$.

we argue that if S is large enough compared to some property of \mathcal{H} , then with high probability all $h \in \mathcal{H}$ have their training error close to their true error, so that if we find a hypothesis whose training error is low, we can be confident its true error will be low as well.

Before giving our first result of this form, we note that it will often be convenient to associate each hypotheses with its $\{-1, 1\}$ -valued indicator function

$$h(x) = \begin{cases} 1 & x \in h \\ -1 & x \notin h \end{cases}$$

In this notation the true error of h is $err_D(h) = \operatorname{Prob}_{x \sim D}[h(x) \neq c^*(x)]$ and the training error is $err_S(h) = \operatorname{Prob}_{x \sim S}[h(x) \neq c^*(x)]$.

6.2 Overfitting and Uniform Convergence

We now present two results that explain how one can guard against overfitting. Given a class of hypotheses \mathcal{H} , the first result states that for any given ϵ greater than zero, so long as the training data set is large compared to $\frac{1}{\epsilon} \ln(|\mathcal{H}|)$, it is unlikely any hypothesis $h \in \mathcal{H}$ will have zero training error but have true error greater than ϵ . This means that with high probability, any hypothesis that our algorithms finds that agrees with the target hypothesis on the training data will have low true error. The second result states that if the training data set is large compared to $\frac{1}{\epsilon^2} \ln(|\mathcal{H}|)$, then it is unlikely that the training error and true error will differ by more than ϵ for any hypothesis in \mathcal{H} . This means that if we find an hypothesis in \mathcal{H} whose training error is low, we can be confident its true error will be low as well, even if its training error is not zero.

The basic idea is the following. If we consider some h with large true error, and we select an element $x \in \mathcal{X}$ at random according to D, there is a reasonable chance that x will belong to the symmetric difference $h \triangle c^*$. If we select a large enough training sample S with each point drawn independently from \mathcal{X} according to D, the chance that S is completely disjoint from $h \triangle c^*$ will be incredibly small. This is just for a single hypothesis h but we can now apply the union bound over all $h \in \mathcal{H}$ of large true error, when \mathcal{H} is finite. We formalize this below.

Theorem 6.1 Let \mathcal{H} be an hypothesis class and let ϵ and δ be greater than zero. If a training set S of size

$$n \ge \frac{1}{\epsilon} (\ln |\mathcal{H}| + \ln(1/\delta)),$$

is drawn from distribution D, then with probability greater than or equal to $1 - \delta$ every hin \mathcal{H} with with true error $\operatorname{err}_D(h) \geq \epsilon$ has training error $\operatorname{err}_S(h) > 0$. Equivalently, with probability greater than or equal to $1 - \delta$, every $h \in \mathcal{H}$ with training error zero has true error less than ϵ .

Proof: Let h_1, h_2, \ldots be the hypotheses in \mathcal{H} with true error greater than or equal to ϵ . These are the hypotheses that we don't want to output. Consider drawing the sample S

	Not spam							Spam								
x_1	x_2	x_3	x_4	x_5	x_6	x_7	x_8	x_9	x_{10}	x_{11}	x_{12}	x_{13}	x_{14}	x_{15}	x_{16}	emails
					\downarrow				\downarrow		\downarrow					
0	0	0	0	0	0	0	0	1	1	1	1	1	1	1	1	target concept
					\uparrow				\uparrow		\uparrow					
0	1	0	0	0	Ò	1	0	1	1	1	Ó	1	0	1	1	hypothesis h_i
					\uparrow				\uparrow		\uparrow					

Figure 6.1: The hypothesis h_i disagrees with the truth in one quarter of the emails. Thus with a training set |S|, the probability that the hypothesis will survive is $(1 - 0.25)^{|S|}$

of size n and let A_i be the event that h_i is consistent with S. Since every h_i has true error greater than or equal to ϵ

$$\operatorname{Prob}(A_i) \leq (1-\epsilon)^n.$$

In other words, if we fix h_i and draw a sample S of size n, the chance that h_i makes no mistakes on S is at most the probability that a coin of bias ϵ comes up tails n times in a row, which is $(1 - \epsilon)^n$. By the union bound over all i we have

$$\operatorname{Prob}\left(\bigcup_{i} A_{i}\right) \leq |\mathcal{H}|(1-\epsilon)^{n}.$$

Using the fact that $(1-\epsilon) \leq e^{-\epsilon}$, the probability that any hypothesis in \mathcal{H} with true error greater than or equal to ϵ has training error zero is at most $|\mathcal{H}|e^{-\epsilon n}$. Replacing n by the sample size bound from the theorem statement, this is at most $|\mathcal{H}|e^{-\ln|\mathcal{H}|-\ln(1/\delta)} = \delta$ as desired.

The conclusion of Theorem 6.1 is sometimes called a "PAC-learning guarantee" since it states that if we can find an $h \in \mathcal{H}$ consistent with the sample, then this h is *P*robably Approximately *C*orrect.

Theorem 6.1 addressed the case where there exists a hypothesis in \mathcal{H} with zero training error. What if the best h_i in \mathcal{H} has 5% error on S? Can we still be confident that its true error is low, say at most 10%? For this, we want an analog of Theorem 6.1 that says for a sufficiently large training set S, every $h_i \in \mathcal{H}$ has training error within $\pm \epsilon$ of the true error with high probability. Such a statement is called *uniform convergence* because we are asking that the training set errors converge to their true errors uniformly over all sets in \mathcal{H} . To see intuitively why such a statement should be true for sufficiently large S and a single hypothesis h_i , consider two strings that differ in 10% of the positions and randomly select a large sample of positions. The number of positions that differ in the sample will be close to 10%.

To prove uniform convergence bounds, we use a tail inequality for sums of independent Bernoulli random variables (i.e., coin tosses). The following is particularly convenient and is a variation on the Chernoff bounds in Section 12.4.11 of the appendix. **Theorem 6.2 (Hoeffding bounds)** Let x_1, x_2, \ldots, x_n be independent $\{0, 1\}$ -valued random variables with probability p that x_i equals one. Let $s = \sum_i x_i$ (equivalently, flip ncoins of bias p and let s be the total number of heads). For any $0 \le \alpha \le 1$,

$$\begin{aligned} \operatorname{Prob}(s/n > p + \alpha) &\leq e^{-2n\alpha^2} \\ \operatorname{Prob}(s/n$$

Theorem 6.2 implies the following uniform convergence analog of Theorem 6.1.

Theorem 6.3 (Uniform convergence) Let \mathcal{H} be a hypothesis class and let ϵ and δ be greater than zero. If a training set S of size

$$n \ge \frac{1}{2\epsilon^2} \big(\ln |\mathcal{H}| + \ln(2/\delta) \big),$$

is drawn from distribution D, then with probability greater than or equal to $1 - \delta$, every h in \mathcal{H} satisfies $|err_S(h) - err_D(h)| \leq \epsilon$.

Proof: First, fix some $h \in \mathcal{H}$ and let x_j be the indicator random variable for the event that h makes a mistake on the j^{th} example in S. The x_j are independent $\{0, 1\}$ random variables and the probability that x_i equals 1 is the true error of h, and the fraction of the x_j 's equal to 1 is exactly the training error of h. Therefore, Hoeffding bounds guarantee that the probability of the event A_h that $|err_D(h) - err_S(h)| > \epsilon$ is less than or equal to $2e^{-2n\epsilon^2}$. Applying the union bound to the events A_h over all $h \in \mathcal{H}$, the probability that there exists an $h \in \mathcal{H}$ with the difference between true error and empirical error greater than ϵ is less than or equal to $2|\mathcal{H}|e^{-2n\epsilon^2}$. Using the value of n from the theorem statement, the right-hand-side of the above inequality is at most δ as desired.

Theorem 6.3 justifies the approach of optimizing over our training sample S even if we are not able to find a rule of zero training error. If our training set S is sufficiently large, with high probability, good performance on S will translate to good performance on D.

Note that Theorems 6.1 and 6.3 require $|\mathcal{H}|$ to be finite in order to be meaningful. The notion of growth functions and VC-dimension in Section 6.9, extend Theorem 6.3 to certain infinite hypothesis classes.

6.3 Illustrative Examples and Occam's Razor

We now present some examples to illustrate the use of Theorem 6.1 and 6.3 and also use these theorems to give a formal connection to the notion of Occam's razor.

6.3.1 Learning disjunctions

Consider the instance space $\mathcal{X} = \{0, 1\}^d$ and suppose we believe that the target concept can be represented by a *disjunction* (an OR) over features, such as $c^* = \{x | x_1 = 1 \lor x_4 = 1 \lor x$

 $1 \vee x_8 = 1$ }, or more succinctly, $c^* = x_1 \vee x_4 \vee x_8$. For example, if we are trying to predict whether an email message is spam or not, and our features correspond to the presence or absence of different possible indicators of spam-ness, then this would correspond to the belief that there is some subset of these indicators such that every spam email has at least one of them and every non-spam email has none of them. Formally, let \mathcal{H} denote the class of disjunctions, and notice that $|\mathcal{H}| = 2^d$. So, by Theorem 6.1, it suffices to find a consistent disjunction over a sample S of size

$$|S| = \frac{1}{\epsilon} \left(d\ln(2) + \ln(1/\delta) \right).$$

How can we efficiently find a consistent disjunction when one exists? Here is a simple algorithm.

Simple Disjunction Learner: Given sample S, discard all features that are set to 1 in any negative example in S. Output the concept h that is the OR of all features that remain.

Lemma 6.4 The Simple Disjunction Learner produces a disjunction h that is consistent with the sample S (i.e., with $err_S(h) = 0$) whenever the target concept is indeed a disjunction.

Proof: Suppose target concept c^* is a disjunction. Then for any x_i that is listed in c^* , x_i will not be set to 1 in any negative example by definition of an OR. Therefore, h will include x_i as well. Since h contains all variables listed in c^* , this ensures that h will correctly predict positive on all positive examples in S. Furthermore, h will correctly predict negative on all negative examples in S since by design all features set to 1 in any negative examples in S.

Thus, combining Lemma 6.4 with Theorem 6.1, we have an efficient algorithm for PAC-learning the class of disjunctions.

6.3.2 Occam's razor

Occam's razor is the notion, stated by William of Occam around AD 1320, that in general one should prefer simpler explanations over more complicated ones.¹⁸ Why should one do this, and can we make a formal claim about why this is a good idea? What if each of us disagrees about precisely which explanations are simpler than others? It turns out we can use Theorem 6.1 to make a mathematical statement of Occam's razor that addresses these issues.

First, what do we mean by a rule being "simple"? Let's assume that each of us has some way of describing rules, using bits (since we are computer scientists). The methods, also called *description languages*, used by each of us may be different, but one fact we can

¹⁸The statement more explicitly was that "Entities should not be multiplied unnecessarily."

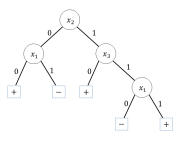


Figure 6.2: A decision tree with three internal nodes and four leaves. This tree corresponds to the Boolean function $\bar{x_1}\bar{x_2} \vee x_1x_2x_3 \vee x_2\bar{x_3}$.

say for certain is that in any given description language, there are at most 2^b rules that can be described using fewer than b bits (because $1 + 2 + 4 + \ldots + 2^{b-1} < 2^b$). Therefore, by setting \mathcal{H} to be the set of all rules that can be described in fewer than b bits and plugging into Theorem 6.1, yields the following:

Theorem 6.5 (Occam's razor) Fix any description language, and consider a training sample S drawn from distribution \mathcal{D} . With probability at least $1 - \delta$, any rule h consistent with S that can be described in this language using fewer than b bits will have $\operatorname{err}_D(h) \leq \epsilon$ for $|S| = \frac{1}{\epsilon} [b \ln(2) + \ln(1/\delta)]$. Equivalently, with probability at least $1 - \delta$, all rules that can be described in fewer than b bits will have $\operatorname{err}_D(h) \leq \frac{b \ln(2) + \ln(1/\delta)}{|S|}$.

For example, using the fact that $\ln(2) < 1$ and ignoring the low-order $\ln(1/\delta)$ term, this means that if the number of bits it takes to write down a rule consistent with the training data is at most 10% of the number of data points in our sample, then we can be confident it will have error at most 10% with respect to \mathcal{D} . What is perhaps surprising about this theorem is that it means that we can each have different ways of describing rules and yet all use Occam's razor. Note that the theorem does not say that complicated rules are necessarily bad, or even that given two rules consistent with the data that the complicated rule is necessarily worse. What it does say is that Occam's razor is a good policy in that simple rules are unlikely to fool us since there are just not that many simple rules.

6.3.3 Application: learning decision trees

One popular practical method for machine learning is to learn a *decision tree*; see Figure 6.2. While finding the smallest decision tree that fits a given training sample S is NP-hard, there are a number of heuristics that are used in practice.¹⁹ Suppose we run such a heuristic on a training set S and it outputs a tree with k nodes. Such a tree can be

¹⁹For instance, one popular heuristic, called ID3, selects the feature to put inside any given node v by choosing the feature of largest *information gain*, a measure of how much it is directly improving prediction. Formally, using S_v to denote the set of examples in S that reach node v, and supposing that feature x_i partitions S_v into S_v^0 and S_v^1 (the examples in S_v with $x_i = 0$ and $x_i = 1$, respectively), the

described using $O(k \log d)$ bits: $\log_2(d)$ bits to give the index of the feature in the root, O(1) bits to indicate for each child if it is a leaf and if so what label it should have, and then $O(k_L \log d)$ and $O(k_R \log d)$ bits respectively to describe the left and right subtrees, where k_L is the number of nodes in the left subtree and k_R is the number of nodes in the right subtree. So, by Theorem 6.5, we can be confident the true error is low if we can produce a consistent tree with fewer than $\epsilon |S|/\log(d)$ nodes.

6.4 Regularization: penalizing complexity

Theorems 6.3 and 6.5 suggest the following idea. Suppose that there is no simple rule that is perfectly consistent with the training data, but we notice there are very simple rules with training error 20%, say, and then some more complex rules with training error 10%, and so on. In this case, perhaps we should optimize some combination of training error and simplicity. This is the notion of *regularization*, also called *complexity penalization*.

Specifically, a *regularizer* is a penalty term that penalizes more complex hypotheses. Given our theorems so far, a natural measure of complexity of a hypothesis is the number of bits we need to write it down.²⁰ Consider now fixing some description language, and let \mathcal{H}_i denote those hypotheses that can be described in *i* bits in this language, so $|\mathcal{H}_i| \leq 2^i$. Let $\delta_i = \delta/2^i$. Rearranging the bound of Theorem 6.3, we know that with probability at least $1 - \delta_i$, all $h \in \mathcal{H}_i$ satisfy $err_D(h) \leq err_S(h) + \sqrt{\frac{\ln(|\mathcal{H}_i|) + \ln(2/\delta_i)}{2|S|}}$. Now, applying the union bound over all *i*, using the fact that $\delta_1 + \delta_2 + \delta_3 + \ldots = \delta$, and also the fact that $\ln(|\mathcal{H}_i|) + \ln(2/\delta_i) \leq i \ln(4) + \ln(2/\delta)$, gives the following corollary.

Corollary 6.6 Fix any description language, and consider a training sample S drawn from distribution \mathcal{D} . With probability greater than or equal to $1 - \delta$, all hypotheses h satisfy

$$err_D(h) \leq err_S(h) + \sqrt{\frac{\operatorname{size}(h)\ln(4) + \ln(2/\delta)}{2|S|}}$$

where size(h) denotes the number of bits needed to describe h in the given language.

Corollary 6.6 gives us the tradeoff we were looking for. It tells us that rather than searching for a rule of low training error, we instead may want to search for a rule with a low right-hand-side in the displayed formula. If we can find one for which this quantity is small, we can be confident true error will be low as well.

information gain of x_i is defined as: $Ent(S_v) - \left[\frac{|S_v^0|}{|S_v|}Ent(S_v^0) + \frac{|S_v^1|}{|S_v|}Ent(S_v^1)\right]$. Here, Ent(S') is the binary entropy of the label proportions in set S'; that is, if a p fraction of the examples in S' are positive, then $Ent(S') = p \log_2(1/p) + (1-p) \log_2(1/(1-p))$, defining $0 \log_2(0) = 0$. This then continues until all leaves are pure—they have only positive or only negative examples.

²⁰Later we will see support vector machines that use a regularizer for linear separators based on the margin of separation of data.

6.5 Online learning and the Perceptron algorithm

So far we have been considering what is often called the *batch learning* scenario. You are given a "batch" of data—the training sample S—and your goal is to use it to produce a hypothesis h that will have low error on new data, under the assumption that both S and the new data are sampled from some fixed distribution D. We now switch to the more challenging *online learning* scenario where we remove the assumption that data is sampled from a fixed probability distribution, or from any probabilistic process at all.

Specifically, the online learning scenario proceeds as follows. At each time t = 1, 2, ...:

- 1. The algorithm is presented with an arbitrary example $x_t \in \mathcal{X}$ and is asked to make a prediction ℓ_t of its label.
- 2. The algorithm is told the true label of the example $c^*(x_t)$ and is charged for a mistake if $c^*(x_t) \neq \ell_t$.

The goal of the learning algorithm is to make as few mistakes as possible in total. For example, consider an email classifier that when a new email message arrives must classify it as "important" or "it can wait". The user then looks at the email and informs the algorithm if it was incorrect. We might not want to model email messages as independent random objects from a fixed probability distribution, because they often are replies to previous emails and build on each other. Thus, the online learning model would be more appropriate than the batch model for this setting.

Intuitively, the online learning model is harder than the batch model because we have removed the requirement that our data consists of independent draws from a fixed probability distribution. Indeed, we will see shortly that any algorithm with good performance in the online model can be converted to an algorithm with good performance in the batch model. Nonetheless, the online model can sometimes be a cleaner model for design and analysis of algorithms.

6.5.1 An example: learning disjunctions

As a simple example, let's revisit the problem of learning disjunctions in the online model. We can solve this problem by starting with a hypothesis $h = x_1 \vee x_2 \vee \ldots \vee x_d$ and using it for prediction. We will maintain the invariant that every variable in the target disjunction is also in our hypothesis, which is clearly true at the start. This ensures that the only mistakes possible are on examples x for which h(x) is positive but $c^*(x)$ is negative. When such a mistake occurs, we simply remove from h any variable set to 1 in x. Since such variables cannot be in the target function (since x was negative), we maintain our invariant and remove at least one variable from h. This implies that the algorithm makes at most d mistakes total on any series of examples consistent with a disjunction. In fact, we can show this bound is tight by showing that no deterministic algorithm can guarantee to make fewer than d mistakes.

Theorem 6.7 For any deterministic algorithm A there exists a sequence of examples σ and disjunction c^* such that A makes at least d mistakes on sequence σ labeled by c^* .

Proof: Let σ be the sequence e_1, e_2, \ldots, e_d where e_j is the example that is zero everywhere except for a 1 in the *j*th position. Imagine running A on sequence σ and telling A it made a mistake on every example; that is, if A predicts positive on e_j we set $c^*(e_j) = -1$ and if A predicts negative on e_j we set $c^*(e_j) = +1$. This target corresponds to the disjunction of all x_j such that A predicted negative on e_j , so it is a legal disjunction. Since A is deterministic, the fact that we constructed c^* by running A is not a problem: it would make the same mistakes if re-run from scratch on the same sequence and same target. Therefore, A makes d mistakes on this σ and c^* .

6.5.2 The Halving algorithm

If we are not concerned with running time, a simple algorithm that guarantees to make at most $\log_2(|\mathcal{H}|)$ mistakes for a target belonging to any given class \mathcal{H} is called the *halving algorithm*. This algorithm simply maintains the version space $\mathcal{V} \subseteq \mathcal{H}$ consisting of all $h \in \mathcal{H}$ consistent with the labels on every example seen so far, and predicts based on majority vote over these functions. Each mistake is guaranteed to reduce the size of the version space \mathcal{V} by at least half (hence the name), thus the total number of mistakes is at most $\log_2(|\mathcal{H}|)$. Note that this can be viewed as the number of bits needed to write a function in \mathcal{H} down.

6.5.3 The Perceptron algorithm

The Perceptron algorithm is an efficient algorithm for learning a linear separator in ddimensional space, with a mistake bound that depends on the margin of separation of the data. Specifically, the assumption is that the target function can be described by a vector \mathbf{w}^* such that for each positive example \mathbf{x} we have $\mathbf{x}^T \mathbf{w}^* \ge 1$ and for each negative example \mathbf{x} we have $\mathbf{x}^T \mathbf{w}^* \le -1$. Note that if we think of the examples \mathbf{x} as points in space, then $\mathbf{x}^T \mathbf{w}^* / |\mathbf{w}^*|$ is the distance of \mathbf{x} to the hyperplane $\mathbf{x}^T \mathbf{w}^* = 0$. Thus, we can view our assumption as stating that there exists a linear separator through the origin with all positive examples on one side, all negative examples on the other side, and all examples at distance at least $\gamma = 1/|\mathbf{w}^*|$ from the separator. This quantity γ is called the margin of separation (see Figure 6.3).

The guarantee of the Perceptron algorithm will be that the total number of mistakes is at most $(R/\gamma)^2$ where $R = \max_t |\mathbf{x}_t|$ over all examples \mathbf{x}_t seen so far. Thus, if there exists a hyperplane through the origin that correctly separates the positive examples from the negative examples by a large margin relative to the radius of the smallest ball enclosing

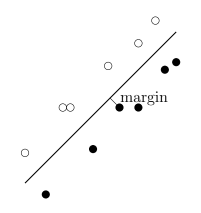


Figure 6.3: Margin of a linear separator.

the data, then the total number of mistakes will be small. The algorithm is very simple and proceeds as follows.

The Perceptron Algorithm: Start with the all-zeroes weight vector w = 0. Then, for t = 1, 2, ... do:

- 1. Given example \mathbf{x}_t , predict $\operatorname{sgn}(\mathbf{x}_t^T \mathbf{w})$.
- 2. If the prediction was a mistake, then update:
 - (a) If \mathbf{x}_t was a positive example, let $\mathbf{w} \leftarrow \mathbf{w} + \mathbf{x}_t$.
 - (b) If \mathbf{x}_t was a negative example, let $\mathbf{w} \leftarrow \mathbf{w} \mathbf{x}_t$.

While simple, the Perceptron algorithm enjoys a strong guarantee on its total number of mistakes.

Theorem 6.8 On any sequence of examples $\mathbf{x}_1, \mathbf{x}_2, \ldots$, if there exists a vector \mathbf{w}^* such that $\mathbf{x}_t^T \mathbf{w}^* \geq 1$ for the positive examples and $\mathbf{x}_t^T \mathbf{w}^* \leq -1$ for the negative examples (i.e., a linear separator of margin $\gamma = 1/|\mathbf{w}^*|$), then the Perceptron algorithm makes at most $R^2|\mathbf{w}^*|^2$ mistakes, where $R = \max_t |\mathbf{x}_t|$.

To get a feel for this bound, notice that if we multiply all entries in all the \mathbf{x}_t by 100, we can divide all entries in \mathbf{w}^* by 100 and it will still satisfy the "if" condition. So the bound is invariant to this kind of scaling, i.e., to what our "units of measurement" are.

Proof of Theorem 6.8: Fix some consistent \mathbf{w}^* . We will keep track of two quantities, $\mathbf{w}^T \mathbf{w}^*$ and $|\mathbf{w}|^2$. First of all, each time we make a mistake, $\mathbf{w}^T \mathbf{w}^*$ increases by at least 1. That is because if \mathbf{x}_t is a positive example, then

$$(\mathbf{w} + \mathbf{x}_t)^T \mathbf{w}^* = \mathbf{w}^T \mathbf{w}^* + \mathbf{x}_t^T \mathbf{w}^* \ge \mathbf{w}^T \mathbf{w}^* + 1,$$

by definition of \mathbf{w}^* . Similarly, if \mathbf{x}_t is a negative example, then

$$(\mathbf{w} - \mathbf{x}_t)^T \mathbf{w}^* = \mathbf{w}^T \mathbf{w}^* - \mathbf{x}_t^T \mathbf{w}^* \ge \mathbf{w}^T \mathbf{w}^* + 1.$$

Next, on each mistake, we claim that $|\mathbf{w}|^2$ increases by at most R^2 . Let us first consider mistakes on positive examples. If we make a mistake on a positive example \mathbf{x}_t then we have

$$(\mathbf{w} + \mathbf{x}_t)^T (\mathbf{w} + \mathbf{x}_t) = |\mathbf{w}|^2 + 2\mathbf{x}_t^T \mathbf{w} + |\mathbf{x}_t|^2 \le |\mathbf{w}|^2 + |\mathbf{x}_t|^2 \le |\mathbf{w}|^2 + R^2,$$

where the middle inequality comes from the fact that we made a mistake, which means that $\mathbf{x}_t^T \mathbf{w} \leq 0$. Similarly, if we make a mistake on a negative example \mathbf{x}_t then we have

$$(\mathbf{w} - \mathbf{x}_t)^T (\mathbf{w} - \mathbf{x}_t) = |\mathbf{w}|^2 - 2\mathbf{x}_t^T \mathbf{w} + |\mathbf{x}_t|^2 \le |\mathbf{w}|^2 + |\mathbf{x}_t|^2 \le |\mathbf{w}|^2 + R^2$$

Note that it is important here that we only update on a mistake.

So, if we make M mistakes, then $\mathbf{w}^T \mathbf{w}^* \ge M$, and $|\mathbf{w}|^2 \le MR^2$, or equivalently, $|\mathbf{w}| \le R\sqrt{M}$. Finally, we use the fact that $\mathbf{w}^T \mathbf{w}^*/|\mathbf{w}^*| \le |\mathbf{w}|$ which is just saying that the projection of \mathbf{w} in the direction of \mathbf{w}^* cannot be larger than the length of \mathbf{w} . This gives us:

$$\begin{array}{rcl} M/|\mathbf{w}^*| &\leq R\sqrt{M} \\ \sqrt{M} &\leq R|\mathbf{w}^*| \\ M &\leq R^2|\mathbf{w}^*|^2 \end{array}$$

as desired.

6.5.4 Extensions: inseparable data and hinge-loss

We assumed above that there existed a perfect \mathbf{w}^* that correctly classified all the examples, e.g., correctly classified all the emails into important versus non-important. This is rarely the case in real-life data. What if even the best \mathbf{w}^* isn't quite perfect? We can see what this does to the above proof: if there is an example that \mathbf{w}^* doesn't correctly classify, then while the second part of the proof still holds, the first part (the dot product of \mathbf{w} with \mathbf{w}^* increasing) breaks down. However, if this doesn't happen too often, and also $\mathbf{x}_t^T \mathbf{w}^*$ is just a "little bit wrong" then we will only make a few more mistakes.

To make this formal, define the *hinge-loss* of \mathbf{w}^* on a positive example \mathbf{x}_t as $\max(0, 1 - \mathbf{x}_t^T \mathbf{w}^*)$. In other words, if $\mathbf{x}_t^T \mathbf{w}^* \ge 1$ as desired then the hinge-loss is zero; else, the hinge-loss is the amount the LHS is less than the RHS.²¹ Similarly, the hinge-loss of \mathbf{w}^* on a negative example \mathbf{x}_t is $\max(0, 1 + \mathbf{x}_t^T \mathbf{w}^*)$. Given a sequence of labeled examples S, define the total hinge-loss $L_{hinge}(\mathbf{w}^*, S)$ as the sum of hinge-losses of \mathbf{w}^* on all examples in S. We now get the following extended theorem.

²¹This is called "hinge-loss" because as a function of $\mathbf{x}_t^T \mathbf{w}^*$ it looks like a hinge.

Theorem 6.9 On any sequence of examples $S = \mathbf{x}_1, \mathbf{x}_2, \ldots$, the Perceptron algorithm makes at most

$$\min_{\mathbf{w}^*} \left(R^2 |\mathbf{w}^*|^2 + 2L_{hinge}(\mathbf{w}^*, S) \right)$$

mistakes, where $R = \max_t |\mathbf{x}_t|$.

Proof: As before, each update of the Perceptron algorithm increases $|\mathbf{w}|^2$ by at most R^2 , so if the algorithm makes M mistakes, we have $|\mathbf{w}|^2 \leq MR^2$.

What we can no longer say is that each update of the algorithm increases $\mathbf{w}^T \mathbf{w}^*$ by at least 1. Instead, on a positive example we are "increasing" $\mathbf{w}^T \mathbf{w}^*$ by $\mathbf{x}_t^T \mathbf{w}^*$ (it could be negative), which is at least $1 - L_{hinge}(\mathbf{w}^*, \mathbf{x}_t)$. Similarly, on a negative example we "increase" $\mathbf{w}^T \mathbf{w}^*$ by $-\mathbf{x}_t^T \mathbf{w}^*$, which is also at least $1 - L_{hinge}(\mathbf{w}^*, \mathbf{x}_t)$. If we sum this up over all mistakes, we get that at the end we have $\mathbf{w}^T \mathbf{w}^* \geq M - L_{hinge}(\mathbf{w}^*, S)$, where we are using here the fact that hinge-loss is never negative so summing over all of S is only larger than summing over the mistakes that \mathbf{w} made.

Finally, we just do some algebra. Let $L = L_{hinge}(\mathbf{w}^*, S)$. So we have:

$$\begin{aligned} \mathbf{w}^{T} \mathbf{w}^{*} / |\mathbf{w}^{*}| &\leq |\mathbf{w}| \\ (\mathbf{w}^{T} \mathbf{w}^{*})^{2} &\leq |\mathbf{w}|^{2} |\mathbf{w}^{*}|^{2} \\ (M - L)^{2} &\leq M R^{2} |\mathbf{w}^{*}|^{2} \\ M^{2} - 2ML + L^{2} &\leq M R^{2} |\mathbf{w}^{*}|^{2} \\ M - 2L + L^{2} / M &\leq R^{2} |\mathbf{w}^{*}|^{2} \\ M &\leq R^{2} |\mathbf{w}^{*}|^{2} + 2L - L^{2} / M \leq R^{2} |\mathbf{w}^{*}|^{2} + 2L \end{aligned}$$

as desired.

6.6 Kernel functions

What if even the best \mathbf{w}^* has high hinge-loss? E.g., perhaps instead of a linear separator decision boundary, the boundary between important emails and unimportant emails looks more like a circle, for example as in Figure 6.4.

A powerful idea for addressing situations like this is to use what are called *kernel* functions, or sometimes the "kernel trick". Here is the idea. Suppose you have a function K, called a "kernel", over pairs of data points such that for some function $\phi : \mathbb{R}^d \to \mathbb{R}^N$, where perhaps $N \gg d$, we have $K(\mathbf{x}, \mathbf{x}') = \phi(\mathbf{x})^T \phi(\mathbf{x}')$. In that case, if we can write the Perceptron algorithm so that it only interacts with the data via dot-products, and then replace every dot-product with an invocation of K, then we can act as if we had performed the function ϕ explicitly without having to actually compute ϕ .

For example, consider $K(\mathbf{x}, \mathbf{x}') = (1 + \mathbf{x}^T \mathbf{x}')^k$ for some integer $k \ge 1$. It turns out this corresponds to a mapping ϕ into a space of dimension $N \approx d^k$. For example, in the case

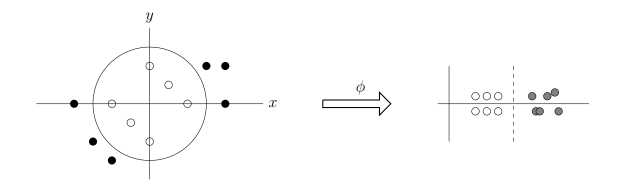


Figure 6.4: Data that is not linearly separable in the input space \mathbb{R}^2 but that is linearly separable in the " ϕ -space," $\phi(\mathbf{x}) = (1, \sqrt{2}x_1, \sqrt{2}x_2, x_1^2, \sqrt{2}x_1x_2, x_2^2)$, corresponding to the kernel function $K(\mathbf{x}^t \mathbf{y}) = (1 + x_1x_2 + y_1y_2)^2$.

d = 2, k = 2 we have (using x_i to denote the *i*th coordinate of **x**):

$$K(\mathbf{x}, \mathbf{x}') = (1 + x_1 x_1' + x_2 x_2')^2$$

= 1 + 2x_1 x_1' + 2x_2 x_2' + x_1^2 x_1'^2 + 2x_1 x_2 x_1' x_2' + x_2^2 x_2'^2
= $\phi(\mathbf{x})^T \phi(\mathbf{x}')$

for $\phi(\mathbf{x}) = (1, \sqrt{2}x_1, \sqrt{2}x_2, x_1^2, \sqrt{2}x_1x_2, x_2^2)$. Notice also that a linear separator in this space could correspond to a more complicated decision boundary such as an ellipse in the original space. For instance, the hyperplane $\phi(\mathbf{x})^T \mathbf{w}^* = 0$ for $\mathbf{w}^* = (-4, 0, 0, 1, 0, 1)$ corresponds to the circle $x_1^2 + x_2^2 = 4$ in the original space, such as in Figure 6.4.

The point of this is that if in the higher-dimensional " ϕ -space" there is a **w**^{*} such that the bound of Theorem 6.9 is small, then the algorithm will perform well and make few mistakes. But the nice thing is we didn't have to computationally perform the mapping ϕ !

So, how can we view the Perceptron algorithm as only interacting with data via dotproducts? Notice that **w** is always a linear combination of data points. For example, if we made mistakes on the first, second and fifth examples, and these examples were positive, positive, and negative respectively, we would have $\mathbf{w} = \mathbf{x}_1 + \mathbf{x}_2 - \mathbf{x}_5$. So, if we keep track of **w** this way, then to predict on a new example \mathbf{x}_t , we can write $\mathbf{x}_t^T \mathbf{w} = \mathbf{x}_t^T \mathbf{x}_1 + \mathbf{x}_t^T \mathbf{x}_2 - \mathbf{x}_t^T \mathbf{x}_5$. So if we just replace each of these dot-products with "K", we are running the algorithm as if we had explicitly performed the ϕ mapping. This is called "kernelizing" the algorithm.

Many different pairwise functions on examples are legal kernel functions. One easy way to create a kernel function is by combining other kernel functions together, via the following theorem.

Theorem 6.10 Suppose K_1 and K_2 are kernel functions. Then

- 1. For any constant $c \ge 0$, cK_1 is a legal kernel. In fact, for any scalar function f, the function $K_3(\mathbf{x}, \mathbf{x}') = f(\mathbf{x})f(\mathbf{x}')K_1(\mathbf{x}, \mathbf{x}')$ is a legal kernel.
- 2. The sum $K_1 + K_2$, is a legal kernel.
- 3. The product, K_1K_2 , is a legal kernel.

You will prove Theorem 6.10 in Exercise 6.9. Notice that this immediately implies that the function $K(\mathbf{x}, \mathbf{x}') = (1 + \mathbf{x}^T \mathbf{x}')^k$ is a legal kernel by using the fact that $K_1(\mathbf{x}, \mathbf{x}') = 1$ is a legal kernel, $K_2(\mathbf{x}, \mathbf{x}') = \mathbf{x}^T \mathbf{x}'$ is a legal kernel, then adding them, and then multiplying that by itself k times. Another popular kernel is the Gaussian kernel, defined as:

$$K(\mathbf{x}, \mathbf{x}') = e^{-c|\mathbf{x}-\mathbf{x}'|^2}.$$

If we think of a kernel as a measure of similarity, then this kernel defines the similarity between two data objects as a quantity that decreases exponentially with the squared distance between them. The Gaussian kernel can be shown to be a true kernel function by first writing it as $f(\mathbf{x})f(\mathbf{x}')e^{2c\mathbf{x}^T\mathbf{x}'}$ for $f(\mathbf{x}) = e^{-c|\mathbf{x}|^2}$ and then taking the Taylor expansion of $e^{2c\mathbf{x}^T\mathbf{x}'}$, applying the rules in Theorem 6.10. Technically, this last step requires considering countably infinitely many applications of the rules and allowing for infinite-dimensional vector spaces.

6.7 Online to Batch Conversion

Suppose we have an online algorithm with a good mistake bound, such as the Perceptron algorithm. Can we use it to get a guarantee in the distributional (batch) learning setting? Intuitively, the answer should be yes since the online setting is only harder. Indeed, this intuition is correct. We present here two natural approaches for such online to batch conversion.

Conversion procedure 1: Random Stopping. Suppose we have an online algorithm \mathcal{A} with mistake-bound M. Say we run the algorithm in a single pass on a sample S of size M/ϵ . Let X_t be the indicator random variable for the event that \mathcal{A} makes a mistake on the tth example. Since $\sum_{t=1}^{|S|} X_t \leq M$ for any set S, we certainly have that $\mathbf{E}[\sum_{t=1}^{|S|} X_t] \leq M$ where the expectation is taken over the random draw of S from $\mathcal{D}^{|S|}$. By linearity of expectation, and dividing both sides by |S| we therefore have:

$$\frac{1}{|S|} \sum_{t=1}^{|S|} \mathbf{E}[X_t] \leq M/|S| = \epsilon.$$
(6.1)

Let h_t denote the hypothesis used by algorithm \mathcal{A} to predict on the *t*th example. Since the *t*th example was randomly drawn from \mathcal{D} , we have $\mathbf{E}[err_{\mathcal{D}}(h_t)] = \mathbf{E}[X_t]$. This means that if we choose *t* at random from 1 to |S|, i.e., stop the algorithm at a random time, the expected error of the resulting prediction rule, taken over the randomness in the draw of *S* and the choice of *t*, is at most ϵ as given by equation (6.1). Thus we have: **Theorem 6.11 (Online to Batch via Random Stopping)** If an online algorithm \mathcal{A} with mistake-bound M is run on a sample S of size M/ϵ and stopped at a random time between 1 and |S|, the expected error of the hypothesis h produced satisfies $\mathbf{E}[err_{\mathcal{D}}(h)] \leq \epsilon$.

Conversion procedure 2: Controlled Testing. A second natural approach to using an online learning algorithm \mathcal{A} in the distributional setting is to just run a series of controlled tests. Specifically, suppose that the initial hypothesis produced by algorithm \mathcal{A} is h_1 . Define $\delta_i = \delta/(i+2)^2$ so we have $\sum_{i=0}^{\infty} \delta_i = (\frac{\pi^2}{6} - 1)\delta \leq \delta$. We draw a set of $n_1 = \frac{1}{\epsilon} \log(\frac{1}{\delta_1})$ random examples and test to see whether h_1 gets all of them correct. Note that if $err_D(h_1) \geq \epsilon$ then the chance h_1 would get them all correct is at most $(1-\epsilon)^{n_1} \leq \delta_1$. So, if h_1 indeed gets them all correct, we output h_1 as our hypothesis and halt. If not, we choose some example x_1 in the sample on which h_1 made a mistake and give it to algorithm \mathcal{A} . Algorithm \mathcal{A} then produces some new hypothesis h_2 and we again repeat, testing h_2 on a fresh set of $n_2 = \frac{1}{\epsilon} \log(\frac{1}{\delta_2})$ random examples, and so on.

In general, given h_t we draw a fresh set of $n_t = \frac{1}{\epsilon} \log(\frac{1}{\delta_t})$ random examples and test to see whether h_t gets all of them correct. If so, we output h_t and halt; if not, we choose some x_t on which $h_t(x_t)$ was incorrect and give it to algorithm \mathcal{A} . By choice of n_t , if h_t had error rate ϵ or larger, the chance we would mistakenly output it is at most δ_t . By choice of the values δ_t , the chance we *ever* halt with a hypothesis of error ϵ or larger is at most $\delta_1 + \delta_2 + \ldots \leq \delta$. Thus, we have the following theorem.

Theorem 6.12 (Online to Batch via Controlled Testing) Let \mathcal{A} be an online learning algorithm with mistake-bound M. Then this procedure will halt after $O(\frac{M}{\epsilon} \log(\frac{M}{\delta}))$ examples and with probability at least $1 - \delta$ will produce a hypothesis of error at most ϵ .

Note that in this conversion we cannot re-use our samples: since the hypothesis h_t depends on the previous data, we need to draw a fresh set of n_t examples to use for testing it.

6.8 Support-Vector Machines

In a batch setting, rather than running the Perceptron algorithm and adapting it via one of the methods above, another natural idea would be just to solve for the vector \mathbf{w} that minimizes the right-hand-side in Theorem 6.9 on the given dataset S. This turns out to have good guarantees as well, though they are beyond the scope of this book. In fact, this is the Support Vector Machine (SVM) algorithm. Specifically, SVMs solve the following convex optimization problem over a sample $S = {\mathbf{x}_1, \mathbf{x}_2, \ldots, \mathbf{x}_n}$ where c is a constant that is determined empirically.

$$\begin{array}{ll} \text{minimize} & c |\mathbf{w}|^2 + \sum_i s_i \\ \text{subject to} & \mathbf{w} \cdot \mathbf{x}_i \geq 1 - s_i \text{ for all positive examples } \mathbf{x}_i \\ & \mathbf{w} \cdot \mathbf{x}_i \leq -1 + s_i \text{ for all negative examples } \mathbf{x}_i \\ & s_i \geq 0 \text{ for all } i. \end{array}$$

Notice that the sum of slack variables is the total hinge loss of \mathbf{w} . So, this convex optimization is minimizing a weighted sum of $1/\gamma^2$, where γ is the margin, and the total hinge loss. If we were to add the constraint that all $s_i = 0$ then this would be solving for the maximum margin linear separator for the data. However, in practice, optimizing a weighted combination generally performs better.

6.9 VC-Dimension

In Section 6.2 we presented several theorems showing that so long as the training set S is large compared to $\frac{1}{\epsilon} \log(|\mathcal{H}|)$, we can be confident that every $h \in \mathcal{H}$ with $err_D(h) \geq \epsilon$ will have $err_S(h) > 0$, and if S is large compared to $\frac{1}{\epsilon^2} \log(|\mathcal{H}|)$, then we can be confident that every $h \in \mathcal{H}$ will have $|err_D(h) - err_S(h)| \leq \epsilon$. In essence, these results used $\log(|\mathcal{H}|)$ as a measure of complexity of class \mathcal{H} . VC-dimension is a different, tighter measure of complexity for a concept class, and as we will see, is also sufficient to yield confidence bounds. For any class \mathcal{H} , VCdim $(\mathcal{H}) \leq \log_2(|\mathcal{H}|)$ but it can also be quite a bit smaller. Let's introduce and motivate it through an example.

Consider a database consisting of the salary and age for a random sample of the adult population in the United States. Suppose we are interested in using the database to answer questions of the form: "what fraction of the adult population in the United States has age between 35 and 45 and salary between \$50,000 and \$70,000?" That is, we are interested in queries that ask about the fraction of the adult population within some axisparallel rectangle. What we can do is calculate the fraction of the database satisfying this condition and return this as our answer. This brings up the following question: How large does our database need to be so that with probability greater than or equal to $1 - \delta$, our answer will be within $\pm \epsilon$ of the truth for *every* possible rectangle query of this form?

If we assume our values are discretized such as 100 possible ages and 1,000 possible salaries, then there are at most $(100 \times 1,000)^2 = 10^{10}$ possible rectangles. This means we can apply Theorem 6.3 with $|\mathcal{H}| \leq 10^{10}$. Specifically, we can think of the target concept c^* as the empty set so that $err_S(h)$ is exactly the fraction of the sample inside rectangle h and $err_D(h)$ is exactly the fraction of the whole population inside h.²² This would tell us that a sample size of $\frac{1}{2\epsilon^2}(10 \ln 10 + \ln(2/\delta))$ would be sufficient.

However, what if we do not wish to discretize our concept class? Another approach would be to say that if there are only N adults total in the United States, then there are at most N^4 rectangles that are truly different with respect to D and so we could use $|\mathcal{H}| \leq N^4$. Still, this suggests that S needs to grow with N, albeit logarithmically, and one might wonder if that is really necessary. VC-dimension, and the notion of the *growth*

²²Technically D is the uniform distribution over the adult population of the United States, and we want to think of S as an independent identical distributed sample from this D.

function of concept class \mathcal{H} , will give us a way to avoid such discretization and avoid any dependence on the size of the support of the underlying distribution D.

6.9.1 Definitions and Key Theorems

Definition 6.1 Given a set S of examples and a concept class \mathcal{H} , we say that S is **shattered** by \mathcal{H} if for every $A \subseteq S$ there exists some $h \in \mathcal{H}$ that labels all examples in A as positive and all examples in $S \setminus A$ as negative.

Definition 6.2 The VC-dimension of \mathcal{H} is the size of the largest set shattered by \mathcal{H} .

For example, there exist sets of four points that can be shattered by rectangles with axis-parallel edges, e.g., four points at the vertices of a diamond (see Figure 6.5). Given such a set S, for any $A \subseteq S$, there exists a rectangle with the points in A inside the rectangle and the points in $S \setminus A$ outside the rectangle. However, rectangles with axis-parallel edges cannot shatter any set of five points. To see this, assume for contradiction that there is a set of five points shattered by the family of axis-parallel rectangles. Find the minimum enclosing rectangle for the five points. For each edge there is at least one point that has stopped its movement. Identify one such point for each edge. The same point may be identified as stopping two edges if it is at a corner of the minimum enclosing rectangle there edge. If two or more points have stopped an edge, designate only one as having stopped the edge. Now, at most four points have been designated. Any rectangle enclosing the designated points must include the undesignated points. Thus, the subset of designated points cannot be expressed as the intersection of a rectangle with the five points. Therefore, the VC-dimension of axis-parallel rectangles is four.

We now need one more definition, which is the growth function of a concept class \mathcal{H} .

Definition 6.3 Given a set S of examples and a concept class \mathcal{H} , let $\mathcal{H}[S] = \{h \cap S : h \in \mathcal{H}\}$. That is, $\mathcal{H}[S]$ is the concept class \mathcal{H} restricted to the set of points S. For integer n and class \mathcal{H} , let $\mathcal{H}[n] = \max_{|S|=n} |\mathcal{H}[S]|$; this is called the **growth function** of \mathcal{H} .

For example, we could have defined shattering by saying that S is shattered by \mathcal{H} if $|\mathcal{H}[S]| = 2^{|S|}$, and then the VC-dimension of \mathcal{H} is the largest n such that $\mathcal{H}[n] = 2^n$. Notice also that for axis-parallel rectangles, $\mathcal{H}[n] = O(n^4)$. The growth function of a class is sometimes called the shatter function or shatter coefficient.

What connects these to learnability are the following three remarkable theorems. The first two are analogs of Theorem 6.1 and Theorem 6.3 respectively, showing that one can replace $|\mathcal{H}|$ with its growth function. This is like replacing the number of concepts in \mathcal{H} with the number of concepts "after the fact", i.e., after S is drawn, and is subtle because we cannot just use a union bound after we have already drawn our set S. The third theorem relates the growth function of a class to its VC-dimension. We now present the theorems, give examples of VC-dimension and growth function of various concept classes, and then prove the theorems.

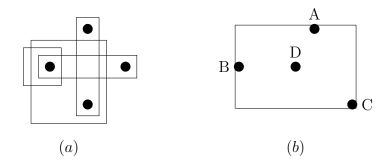


Figure 6.5: (a) shows a set of four points that can be shattered by rectangles along with some of the rectangles that shatter the set. Not every set of four points can be shattered as seen in (b). Any rectangle containing points A, B, and C must contain D. No set of five points can be shattered by rectangles with axis-parallel edges. No set of three collinear points can be shattered, since any rectangle that contains the two end points must also contain the middle point. More generally, since rectangles are convex, a set with one point inside the convex hull of the others cannot be shattered.

Theorem 6.13 (Growth function sample bound) For any class \mathcal{H} and distribution \mathcal{D} , if a training sample S is drawn from \mathcal{D} of size

$$n \geq \frac{2}{\epsilon} [\log_2(2\mathcal{H}[2n]) + \log_2(1/\delta)]$$

then with probability $\geq 1 - \delta$, every $h \in \mathcal{H}$ with $err_{\mathcal{D}}(h) \geq \epsilon$ has $err_{S}(h) > 0$ (equivalently, every $h \in \mathcal{H}$ with $err_{S}(h) = 0$ has $err_{\mathcal{D}}(h) < \epsilon$).

Theorem 6.14 (Growth function uniform convergence) For any class \mathcal{H} and distribution \mathcal{D} , if a training sample S is drawn from \mathcal{D} of size

$$n \geq \frac{8}{\epsilon^2} [\ln(2\mathcal{H}[2n]) + \ln(1/\delta)]$$

then with probability $\geq 1 - \delta$, every $h \in \mathcal{H}$ will have $|err_S(h) - err_D(h)| \leq \epsilon$.

Theorem 6.15 (Sauer's lemma) If $\operatorname{VCdim}(\mathcal{H}) = d$ then $\mathcal{H}[n] \leq \sum_{i=0}^{d} {n \choose i} \leq (\frac{en}{d})^{d}$.

Notice that Sauer's lemma was fairly tight in the case of axis-parallel rectangles, though in some cases it can be a bit loose. E.g., we will see that for linear separators in the plane, their VC-dimension is 3 but $\mathcal{H}[n] = O(n^2)$. An interesting feature about Sauer's lemma is that it implies the growth function switches from taking the form 2^n to taking the form $n^{\text{VCdim}(\mathcal{H})}$ when n reaches the VC-dimension of the class \mathcal{H} .

Putting Theorems 6.13 and 6.15 together, with a little algebra we get the following corollary (a similar corollary results by combining Theorems 6.14 and 6.15):

Corollary 6.16 (VC-dimension sample bound) For any class \mathcal{H} and distribution \mathcal{D} , a training sample S of size

$$O\left(\frac{1}{\epsilon}[\operatorname{VCdim}(\mathcal{H})\log(1/\epsilon) + \log(1/\delta)]\right)$$

is sufficient to ensure that with probability $\geq 1 - \delta$, every $h \in \mathcal{H}$ with $err_{\mathcal{D}}(h) \geq \epsilon$ has $err_{S}(h) > 0$ (equivalently, every $h \in \mathcal{H}$ with $err_{S}(h) = 0$ has $err_{\mathcal{D}}(h) < \epsilon$).

For any class \mathcal{H} , VCdim $(\mathcal{H}) \leq \log_2(|\mathcal{H}|)$ since \mathcal{H} must have at least 2^k concepts in order to shatter k points. Thus Corollary 6.16 is never too much worse than Theorem 6.1 and can be much better.

6.9.2 Examples: VC-Dimension and Growth Function

Rectangles with axis-parallel edges

As we saw above, the class of axis-parallel rectangles in the plane has VC-dimension 4 and growth function $C[n] = O(n^4)$.

Intervals of the reals

Intervals on the real line can shatter any set of two points but no set of three points since the subset of the first and last points cannot be isolated. Thus, the VC-dimension of intervals is two. Also, $C[n] = O(n^2)$ since we have $O(n^2)$ choices for the left and right endpoints.

Pairs of intervals of the reals

Consider the family of pairs of intervals, where a pair of intervals is viewed as the set of points that are in at least one of the intervals, in other words, their set union. There exists a set of size four that can be shattered but no set of size five since the subset of first, third, and last point cannot be isolated. Thus, the VC-dimension of pairs of intervals is four. Also we have $C[n] = O(n^4)$.

Convex polygons

Consider the set system of all convex polygons in the plane. For any positive integer n, place n points on the unit circle. Any subset of the points are the vertices of a convex polygon. Clearly that polygon will not contain any of the points not in the subset. This shows that convex polygons can shatter arbitrarily large sets, so the VC-dimension is infinite. Notice that this also implies that $C[n] = 2^n$.

Half spaces in *d*-dimensions

Define a half space to be the set of all points on one side of a hyper plane, i.e., a set of the form $\{\mathbf{x} | \mathbf{w}^T \mathbf{x} \ge w_0\}$. The VC-dimension of half spaces in *d*-dimensions is d + 1.

There exists a set of size d + 1 that can be shattered by half spaces. Select the d unit-coordinate vectors plus the origin to be the d + 1 points. Suppose A is any subset of these d + 1 points. Without loss of generality assume that the origin is in A. Take a 0-1 vector \mathbf{w} which has 1's precisely in the coordinates corresponding to vectors not in A. Clearly A lies in the half-space $\mathbf{w}^T \mathbf{x} \leq 0$ and the complement of A lies in the complementary half-space.

We now show that no set of d + 2 points in *d*-dimensions can be shattered by linear separators. This is done by proving that any set of d+2 points can be partitioned into two disjoint subsets *A* and *B* of points whose convex hulls intersect. This establishes the claim since any linear separator with *A* on one side must have its entire convex hull on that side,²³ so it is not possible to have a linear separator with *A* on one side and *B* on the other.

Let convex(S) denote the convex hull of point set S.

Theorem 6.17 (Radon): Any set $S \subseteq \mathbb{R}^d$ with $|S| \ge d+2$, can be partitioned into two disjoint subsets A and B such that $convex(A) \cap convex(B) \ne \phi$.

Proof: Without loss of generality, assume |S| = d+2. Form a $d \times (d+2)$ matrix with one column for each point of S. Call the matrix A. Add an extra row of all 1's to construct a $(d+1) \times (d+2)$ matrix B. Clearly the rank of this matrix is at most d+1 and the columns are linearly dependent. Say $\mathbf{x} = (x_1, x_2, \ldots, x_{d+2})$ is a nonzero vector with $B\mathbf{x} = 0$. Reorder the columns so that $x_1, x_2, \ldots, x_s \ge 0$ and $x_{s+1}, x_{s+2}, \ldots, x_{d+2} < 0$. Normalize \mathbf{x} so $\sum_{i=1}^{s} |x_i| = 1$. Let $\mathbf{b_i}$ (respectively $\mathbf{a_i}$) be the i^{th} column of B (respectively A). Then, $\sum_{i=1}^{s} |x_i| \mathbf{b_i} = \sum_{i=s+1}^{d+2} |x_i| \mathbf{b_i}$ from which it follows that $\sum_{i=1}^{s} |x_i| \mathbf{a_i} = \sum_{i=s+1}^{d+2} |x_i| \mathbf{a_i}$ and $\sum_{i=1}^{s} |x_i| = \sum_{i=s+1}^{d+2} |x_i|$. Since $\sum_{i=1}^{s} |x_i| = 1$ and $\sum_{i=s+1}^{d+2} |x_i| = 1$ each side of $\sum_{i=1}^{s} |x_i| \mathbf{a_i} = \sum_{i=s+1}^{d+2} |x_i| \mathbf{a_i}$ is a convex combination of columns of A which proves the theorem. Thus, S can be partitioned into two sets, the first consisting of the first s points after the rearrangement and the second consisting of points s + 1 through d + 2. Their convex hulls intersect as required.

Radon's theorem immediately implies that half-spaces in d-dimensions do not shatter any set of d + 2 points.

Spheres in *d*-dimensions

²³If any two points \mathbf{x}_1 and \mathbf{x}_2 lie on the same side of a separator, so must any convex combination: if $\mathbf{w} \cdot \mathbf{x}_1 \ge b$ and $\mathbf{w} \cdot \mathbf{x}_2 \ge b$ then $\mathbf{w} \cdot (a\mathbf{x}_1 + (1-a)\mathbf{x}_2) \ge b$.

A sphere in d-dimensions is a set of points of the form $\{\mathbf{x} \mid |\mathbf{x} - \mathbf{x}_0| \leq r\}$. The VCdimension of spheres is d + 1. It is the same as that of half spaces. First, we prove that no set of d + 2 points can be shattered by spheres. Suppose some set S with d + 2 points can be shattered. Then for any partition A_1 and A_2 of S, there are spheres B_1 and B_2 such that $B_1 \cap S = A_1$ and $B_2 \cap S = A_2$. Now B_1 and B_2 may intersect, but there is no point of S in their intersection. It is easy to see that there is a hyperplane perpendicular to the line joining the centers of the two spheres with all of A_1 on one side and all of A_2 on the other and this implies that half spaces shatter S, a contradiction. Therefore no d+2 points can be shattered by hyperspheres.

It is also not difficult to see that the set of d+1 points consisting of the unit-coordinate vectors and the origin can be shattered by spheres. Suppose A is a subset of the d+1 points. Let a be the number of unit vectors in A. The center \mathbf{a}_0 of our sphere will be the sum of the vectors in A. For every unit vector in A, its distance to this center will be $\sqrt{a-1}$ and for every unit vector outside A, its distance to this center will be $\sqrt{a+1}$. The distance of the origin to the center is \sqrt{a} . Thus, we can choose the radius so that precisely the points in A are in the hypersphere.

Finite sets

The system of finite sets of real numbers can shatter any finite set of real numbers and thus the VC-dimension of finite sets is infinite.

6.9.3 **Proof of Main Theorems**

We begin with a technical lemma. Consider drawing a set S of n examples from \mathcal{D} and let A denote the event that there exists $h \in \mathcal{H}$ with zero training error on S but true error greater than or equal to ϵ . Now draw a second set S' of n examples from \mathcal{D} and let B denote the event that there exists $h \in \mathcal{H}$ with zero error on S but error greater than or equal to $\epsilon/2$ on S'.

Lemma 6.18 Let \mathcal{H} be a concept class over some domain \mathcal{X} and let S and S' be sets of n elements drawn from some distribution \mathcal{D} on \mathcal{X} , where $n \geq 8/\epsilon$. Let A be the event that there exists $h \in \mathcal{H}$ with zero error on S but true error greater than or equal to ϵ . Let B be the event that there exists $h \in \mathcal{H}$ with zero error on S but error greater than or equal to ϵ . In $f(B) \geq Prob(A)/2$.

Proof: Clearly, $\operatorname{Prob}(B) \geq \operatorname{Prob}(A, B) = \operatorname{Prob}(A)\operatorname{Prob}(B|A)$. Consider drawing set S and suppose event A occurs. Let h be in \mathcal{H} with $\operatorname{err}_{\mathcal{D}}(h) \geq \epsilon$ but $\operatorname{err}_{S}(h) = 0$. Now, draw set S'. E(error of h on S') = $\operatorname{err}_{\mathcal{D}}(h) \geq \epsilon$. So, by Chernoff bounds, since $n \geq 8/\epsilon$, $\operatorname{Prob}(\operatorname{err}_{S'}(h) \geq \epsilon/2) \geq 1/2$. Thus, $\operatorname{Prob}(B|A) \geq 1/2$ and $\operatorname{Prob}(B) \geq \operatorname{Prob}(A)/2$ as desired.

We now prove Theorem 6.13, restated here for convenience.

Theorem 6.13 (Growth function sample bound) For any class \mathcal{H} and distribution \mathcal{D} , if a training sample S is drawn from \mathcal{D} of size

$$n \geq \frac{2}{\epsilon} [\log_2(2\mathcal{H}[2n]) + \log_2(1/\delta)]$$

then with probability $\geq 1 - \delta$, every $h \in \mathcal{H}$ with $err_{\mathcal{D}}(h) \geq \epsilon$ has $err_{S}(h) > 0$ (equivalently, every $h \in \mathcal{H}$ with $err_{S}(h) = 0$ has $err_{\mathcal{D}}(h) < \epsilon$).

Proof: Consider drawing a set S of n examples from \mathcal{D} and let A denote the event that there exists $h \in \mathcal{H}$ with true error greater than ϵ but training error zero. Our goal is to prove that $\operatorname{Prob}(A) \leq \delta$.

By Lemma 6.18 it suffices to prove that $\operatorname{Prob}(B) \leq \delta/2$. Consider a third experiment. Draw a set S'' of 2n points from \mathcal{D} and then randomly partition S'' into two sets S and S' of n points each. Let B^* denote the event that there exists $h \in \mathcal{H}$ with $err_S(h) = 0$ but $err_{S'}(h) \geq \epsilon/2$. $\operatorname{Prob}(B^*) = \operatorname{Prob}(B)$ since drawing 2n points from \mathcal{D} and randomly partitioning them into two sets of size n produces the same distribution on (S, S') as does drawing S and S' directly. The advantage of this new experiment is that we can now argue that $\operatorname{Prob}(B^*)$ is low by arguing that for any set S'' of size 2n, $\operatorname{Prob}(B^*|S'')$ is low, with probability now taken over just the random partition of S'' into S and S'. The key point is that since S'' is fixed, there are at most $|\mathcal{H}[S'']| \leq \mathcal{H}[2n]$ events to worry about. Specifically, it suffices to prove that for any fixed $h \in \mathcal{H}[S'']$, the probability over the partition of S'' that h makes zero mistakes on S but more than $\epsilon n/2$ mistakes on S' is at most $\delta/(2\mathcal{H}[2n])$. We can then apply the union bound over $\mathcal{H}[S''] = \{h \cap S'' | h \in \mathcal{H}\}$.

To make the calculations easier, consider the following specific method for partitioning S'' into S and S'. Randomly put the points in S'' into pairs: $(a_1, b_1), (a_2, b_2), \ldots, (a_n, b_n)$. For each index i, flip a fair coin. If heads put a_i into S and b_i into S', else if tails put a_i into S' and b_i into S. Now, fix some partition $h \in \mathcal{H}[S'']$ and consider the probability over these n fair coin flips that h makes zero mistakes on S but more than $\epsilon n/2$ mistakes on S'. First of all, if for any index i, h makes a mistake on both a_i and b_i then the probability is zero (because it cannot possibly make zero mistakes on S). Second, if there are fewer than $\epsilon n/2$ indices i such that h makes a mistake on either a_i or b_i then again the probability is zero because it cannot possibly make more than $\epsilon n/2$ mistakes on S'. So, assume there are $r \geq \epsilon n/2$ indices i such that h makes a mistake on exactly one of a_i or b_i . In this case, the chance that all of those mistakes land in S' is exactly $1/2^r$. This quantity is at most $1/2^{\epsilon n/2} \leq \delta/(2\mathcal{H}[2n])$ as desired for n as given in the theorem statement.

We now prove Theorem 6.14, restated here for convenience.

Theorem 6.14 (Growth function uniform convergence) For any class \mathcal{H} and distribution \mathcal{D} , if a training sample S is drawn from \mathcal{D} of size

$$n \geq \frac{8}{\epsilon^2} [\ln(2\mathcal{H}[2n]) + \ln(1/\delta)]$$

then with probability $\geq 1 - \delta$, every $h \in \mathcal{H}$ will have $|err_S(h) - err_D(h)| \leq \epsilon$.

Proof: This proof is identical to the proof of Theorem 6.13 except B^* is now the event that there exists a set $h \in \mathcal{H}[S'']$ such that the error of h on S differs from the error of h on S' by more than $\epsilon/2$. We again consider the experiment where we randomly put the points in S'' into pairs (a_i, b_i) and then flip a fair coin for each index i, if heads placing a_i into S and b_i into S', else placing a_i into S' and b_i into S. Consider the difference between the number of mistakes h makes on S and the number of mistakes h makes on S' and observe how this difference changes as we flip coins for $i = 1, 2, \ldots, n$. Initially, the difference is zero. If h makes a mistake on both or neither of (a_i, b_i) then the difference does not change. Else, if h makes a mistake on exactly one of a_i or b_i , then with probability 1/2 the difference increases by one and with probability 1/2 the difference decreases by one. If there are $r \leq n$ such pairs, then if we take a random walk of $r \leq n$ steps, what is the probability that we end up more than $\epsilon n/2$ steps away from the origin? This is equivalent to asking: if we flip $r \leq n$ fair coins, what is the probability the number of heads differes from its expectation by more than $\epsilon n/4$. By Hoeffding bounds, this is at most $2e^{-\epsilon^2 n/8}$.

Finally, we prove Sauer's lemma, relating the growth function to the VC-dimension.

Theorem 6.15 (Sauer's lemma) If $\operatorname{VCdim}(\mathcal{H}) = d$ then $\mathcal{H}[n] \leq \sum_{i=0}^{d} {n \choose i} \leq (\frac{en}{d})^{d}$.

Proof: Let $d = \operatorname{VCdim}(\mathcal{H})$. Our goal is to prove for any set S of n points that $|\mathcal{H}[S]| \leq \binom{n}{\leq d}$, where we are defining $\binom{n}{\leq d} = \sum_{i=0}^{d} \binom{n}{i}$; this is the number of distinct ways of choosing d or fewer elements out of n. We will do so by induction on n. As a base case, our theorem is trivially true if $n \leq d$.

As a first step in the proof, notice that:

$$\binom{n}{\leq d} = \binom{n-1}{\leq d} + \binom{n-1}{\leq d-1}$$
(6.2)

because we can partition the ways of choosing d or fewer items into those that do not include the first item (leaving $\leq d$ to be chosen from the remainder) and those that do include the first item (leaving $\leq d - 1$ to be chosen from the remainder).

Now, consider any set S of n points and pick some arbitrary point $x \in S$. By induction, we may assume that $|\mathcal{H}[S \setminus \{x\}]| \leq \binom{n-1}{\leq d}$. So, by equation (6.2) all we need to show is that $|\mathcal{H}[S]| - |\mathcal{H}[S \setminus \{x\}]| \leq \binom{n-1}{\leq d-1}$. Thus, our problem has reduced to analyzing how many *more* partitions there are of S than there are of $S \setminus \{x\}$ using sets in \mathcal{H} .

If $\mathcal{H}[S]$ is larger than $\mathcal{H}[S \setminus \{x\}]$, it is because of pairs of sets in $\mathcal{H}[S]$ that differ only on point x and therefore collapse to the same set when x is removed. For set $h \in \mathcal{H}[S]$ containing point x, define twin(h) = $h \setminus \{x\}$; this may or may not belong to $\mathcal{H}[S]$. Let $\mathcal{T} = \{h \in \mathcal{H}[S] : x \in h \text{ and twin}(h) \in \mathcal{H}[S]\}$. Notice $|\mathcal{H}[S]| - |\mathcal{H}[S \setminus \{x\}]| = |\mathcal{T}|$.

Now, what is the VC-dimension of \mathcal{T} ? If $d' = \operatorname{VCdim}(\mathcal{T})$, this means there is some set R of d' points in $S \setminus \{x\}$ that are shattered by \mathcal{T} . By definition of \mathcal{T} , all $2^{d'}$ subsets of R can be extended to either include x, or not include x and still be a set in $\mathcal{H}[S]$. In other words, $R \cup \{x\}$ is shattered by \mathcal{H} . This means, $d' + 1 \leq d$. Since $\operatorname{VCdim}(\mathcal{T}) \leq d - 1$, by induction we have $|\mathcal{T}| \leq {\binom{n-1}{d-1}}$ as desired.

6.9.4 VC-dimension of combinations of concepts

Often one wants to create concepts out of other concepts. For example, given several linear separators, one could take their intersection to create a convex polytope. Or given several disjunctions, one might want to take their majority vote. We can use Sauer's lemma to show that such combinations do not increase the VC-dimension of the class by too much.

Specifically, given k concepts h_1, h_2, \ldots, h_k and a Booelan function f define the set $comb_f(h_1, \ldots, h_k) = \{x \in \mathcal{X} : f(h_1(x), \ldots, h_k(x)) = 1\}$, where here we are using $h_i(x)$ to denote the indicator for whether or not $x \in h_i$. For example, f might be the AND function to take the intersection of the sets h_i , or f might be the majority-vote function. This can be viewed as a *depth-two neural network*. Given a concept class \mathcal{H} , a Boolean function f, and an integer k, define the new concept class $COMB_{f,k}(\mathcal{H}) = \{comb_f(h_1, \ldots, h_k) : h_i \in \mathcal{H}\}$. We can now use Sauer's lemma to produce the following corollary.

Corollary 6.19 If the concept class \mathcal{H} has VC-dimension d, then for any combination function f, the class $COMB_{f,k}(\mathcal{H})$ has VC-dimension $O(kd \log(kd))$.

Proof: Let *n* be the VC-dimension of $\text{COMB}_{f,k}(\mathcal{H})$, so by definition, there must exist a set *S* of *n* points shattered by $\text{COMB}_{f,k}(\mathcal{H})$. We know by Sauer's lemma that there are at most n^d ways of partitioning the points in *S* using sets in \mathcal{H} . Since each set in $\text{COMB}_{f,k}(\mathcal{H})$ is determined by *k* sets in \mathcal{H} , and there are at most $(n^d)^k = n^{kd}$ different *k*-tuples of such sets, this means there are at most n^{kd} ways of partitioning the points using sets in $\text{COMB}_{f,k}(\mathcal{H})$. Since *S* is shattered, we must have $2^n \leq n^{kd}$, or equivalently $n \leq kd \log_2(n)$. We solve this as follows. First, assuming $n \geq 16$ we have $\log_2(n) \leq \sqrt{n}$ so $kd \log_2(n) \leq kd\sqrt{n}$ which implies that $n \leq (kd)^2$. To get the better bound, plug back into the original inequality. Since $n \leq (kd)^2$, it must be that $\log_2(n) \leq 2 \log_2(kd)$. substituting $\log n \leq 2 \log_2(kd)$ into $n \leq kd \log_2 n$ gives $n \leq 2kd \log_2(kd)$.

This result will be useful for our discussion of Boosting in Section 6.10.

6.9.5 Other measures of complexity

VC-dimension and number of bits needed to describe a set are not the only measures of complexity one can use to derive generalization guarantees. There has been significant work on a variety of measures. One measure called Rademacher complexity measures the extent to which a given concept class \mathcal{H} can fit random noise. Given a set of nexamples $S = \{x_1, \ldots, x_n\}$, the *empirical Rademacher complexity* of \mathcal{H} is defined as $R_S(\mathcal{H}) = \mathbf{E}_{\sigma_1,\ldots,\sigma_n} \max_{h\in\mathcal{H}} \frac{1}{n} \sum_{i=1}^n \sigma_i h(x_i)$, where $\sigma_i \in \{-1,1\}$ are independent random labels with $\operatorname{Prob}[\sigma_i = 1] = \frac{1}{2}$. E.g., if you assign random ± 1 labels to the points in S and the best classifier in \mathcal{H} on average gets error 0.45 then $R_S(\mathcal{H}) = 0.55 - 0.45 = 0.1$. One can prove that with probability greater than or equal to $1 - \delta$, every $h \in \mathcal{H}$ satisfies true error less than or equal to training error plus $R_S(\mathcal{H}) + 3\sqrt{\frac{\ln(2/\delta)}{2n}}$. For more on results such as this, see, e.g., [?].

6.10 Strong and Weak Learning - Boosting

We now describe *boosting*, which is important both as a theoretical result and as a practical and easy-to-use learning method.

A strong learner for a problem is an algorithm that with high probability is able to achieve any desired error rate ϵ using a number of samples that may depend polynomially on $1/\epsilon$. A weak learner for a problem is an algorithm that does just a little bit better than random guessing. It is only required to get with high probability an error rate less than or equal to $\frac{1}{2} - \gamma$ for some $0 < \gamma \leq \frac{1}{2}$. We show here that a weak-learner for a problem that achieves the weak-learning guarantee for any distribution of data can be boosted to a strong learner, using the technique of boosting. At the high level, the idea will be to take our training sample S, and then to run the weak-learner on different data distributions produced by weighting the points in the training sample in different ways. Running the weak learner on these different weightings of the training sample will produce a series of hypotheses h_1, h_2, \ldots , and the idea of our reweighting procedure will be to focus attention on the parts of the sample that previous hypotheses have performed poorly on. At the end we will combine the hypotheses together by a majority vote.

Assume the weak learning algorithm A outputs hypotheses from some class \mathcal{H} . Our boosting algorithm will produce hypotheses that will be majority votes over t_0 hypotheses from \mathcal{H} , for t_0 defined below. This means that we can apply Corollary 6.19 to bound the VC-dimension of the class of hypotheses our boosting algorithm can produce in terms of the VC-dimension of \mathcal{H} . In particular, the class of rules that can be produced by the booster running for t_0 rounds has VC-dimension $O(t_0 \text{VCdim}(\mathcal{H}) \log(t_0 \text{VCdim}(\mathcal{H})))$. This in turn gives a bound on the number of samples needed, via Corollary 6.16, to ensure that high accuracy on the sample will translate to high accuracy on new data.

To make the discussion simpler, we will assume that the weak learning algorithm A, when presented with a weighting of the points in our training sample, always (rather than with high probability) produces a hypothesis that performs slightly better than random guessing with respect to the distribution induced by weighting. Specificially:

Boosting Algorithm

Given a sample S of n labeled examples $\mathbf{x}_1, \ldots, \mathbf{x}_n$, initialize each example \mathbf{x}_i to have a weight $w_i = 1$. Let $\mathbf{w} = (w_1, \ldots, w_n)$.

For $t = 1, 2, ..., t_0$ do

Call the weak learner on the weighted sample (S, \mathbf{w}) , receiving hypothesis h_t .

Multiply the weight of each example that was misclassified by h_t by $\alpha = \frac{\frac{1}{2} + \gamma}{\frac{1}{2} - \gamma}$. Leave the other weights as they are.

End

Output the classifier $MAJ(h_1, \ldots, h_{t_0})$ which takes the majority vote of the hypotheses returned by the weak learner. Assume t_0 is odd so there is no tie.

Figure 6.6: The boosting algorithm

Definition 6.4 (γ **-Weak learner on sample)** A weak learner is an algorithm that given examples, their labels, and a nonnegative real weight w_i on each example \mathbf{x}_i , produces a classifier that correctly labels a subset of examples with total weight at least $(\frac{1}{2} + \gamma) \sum_{i=1}^{n} w_i$.

At the high level, boosting makes use of the intuitive notion that if an example was misclassified, one needs to pay more attention to it. The boosting procedure is in Figure 6.6.

Theorem 6.20 Let A be a γ -weak learner for sample S. Then $t_0 = O(\frac{1}{\gamma^2} \log n)$ is sufficient so that the classifier $MAJ(h_1, \ldots, h_{t_0})$ produced by the boosting procedure has training error zero.

Proof: Suppose *m* is the number of examples the final classifier gets wrong. Each of these *m* examples was misclassified at least $t_0/2$ times so each has weight at least $\alpha^{t_0/2}$. Thus the total weight is at least $m\alpha^{t_0/2}$. On the other hand, at time t+1, only the weights of examples misclassified at time *t* were increased. By the property of weak learning, the total weight of misclassified examples is at most $(\frac{1}{2} - \gamma)$ of the total weight at time *t*. Let weight(*t*) be the total weight at time *t*. Then

weight
$$(t+1) \leq \left(\alpha \left(\frac{1}{2} - \gamma\right) + \left(\frac{1}{2} + \gamma\right)\right) \times \text{weight}(t)$$

= $(1+2\gamma) \times \text{weight}(t).$

Since weight (0) = n, the total weight at the end is at most $n(1+2\gamma)^{t_0}$. Thus

$$m\alpha^{t_0/2} \leq \text{total weight at end} \leq n(1+2\gamma)^{t_0}.$$

Substituting $\alpha = \frac{1/2+\gamma}{1/2-\gamma} = \frac{1+2\gamma}{1-2\gamma}$ and rearranging terms

$$m \leq n(1-2\gamma)^{t_0/2}(1+2\gamma)^{t_0/2} = n[1-4\gamma^2]^{t_0/2}.$$

Using $1 - x \leq e^{-x}$, $m \leq n e^{-2t_0 \gamma^2}$. For $t_0 > \frac{\ln n}{2\gamma^2}$, m < 1, so the number of misclassified items must be zero.

Having completed the proof of the boosting result, here are two interesting observations:

- **Connection to Hoeffding bounds:** The boosting result applies even if our weak learning algorithm is "adversarial", giving us the least helpful classifier possible subject to Definition 6.4. This is why we don't want the α in the boosting algorithm to be too large, otherwise the weak learner could return the negation of the classifier it gave the last time. Suppose that the weak learning algorithm gave a classifier each time that for each example, flipped a coin and produced the correct answer with probability $\frac{1}{2} + \gamma$ and the wrong answer with probability $\frac{1}{2} - \gamma$, so it is a γ -weak learner in expectation. In that case, if we called the weak learner t_0 times, for any fixed \mathbf{x}_i , Hoeffding bounds imply the chance the majority vote of those classifiers is incorrect on \mathbf{x}_i is at most $e^{-2t_0\gamma^2}$. So, the expected total number of mistakes m is at most $ne^{-2t_0\gamma^2}$. What is interesting is that this is the exact bound we get from boosting without the expectation for an adversarial weak-learner.
- A minimax view: Consider a 2-player zero-sum game ²⁴ with one row for each example \mathbf{x}_i and one column for each hypothesis h_j that the weak-learning algorithm might output. If the row player chooses row i and the column player chooses column j, then the column player gets a payoff of one if $h_j(\mathbf{x}_i)$ is correct and gets a payoff of zero if $h_j(\mathbf{x}_i)$ is incorrect. The γ -weak learning assumption implies that for any randomized strategy for the row player (any "mixed strategy" in the language of game theory), there exists a response h_j that gives the column player an expected payoff of at least $\frac{1}{2} + \gamma$. The von Neumann minimax theorem ²⁵ states that this implies there exists a probability distribution on the columns (a mixed strategy for the column player) such that for any \mathbf{x}_i , at least a $\frac{1}{2} + \gamma$ probability mass of the

 $^{^{24}}$ A two person zero sum game consists of a matrix whose columns correspond to moves for Player 1 and whose rows correspond to moves for Player 2. The ij^{th} entry of the matrix is the payoff for Player 1 if Player 1 choose the j^{th} column and Player 2 choose the i^{th} row. Player 2's payoff is the negative of Player1's.

²⁵The von Neumann minimax theorem states that there exists a mixed strategy for each player so that given Player 2's strategy the best payoff possible for Player 1 is the negative of given Player 1's strategy the best possible payoff for Player 2. A mixed strategy is one in which a probability is assigned to every possible move for each situation a player could be in.

columns under this distribution is correct on \mathbf{x}_i . We can think of boosting as a fast way of finding a very simple probability distribution on the columns (just an average over $O(\log n)$ columns, possibly with repetitions) that is nearly as good (for any \mathbf{x}_i , more than half are correct) that moreover works even if our only access to the columns is by running the weak learner and observing its outputs.

We argued above that $t_0 = O(\frac{1}{\gamma^2} \log n)$ rounds of boosting are sufficient to produce a majority-vote rule h that will classify all of S correctly. Using our VC-dimension bounds, this implies that if the weak learner is choosing its hypotheses from concept class \mathcal{H} , then a sample size

$$n = \tilde{O}\left(\frac{1}{\epsilon} \left(\frac{\operatorname{VCdim}(\mathcal{H})}{\gamma^2}\right)\right)$$

is sufficient to conclude that with probability $1 - \delta$ the error is less than or equal to ϵ , where we are using the \tilde{O} notation to hide logarithmic factors. It turns out that running the boosting procedure for larger values of t_0 i.e., continuing past the point where S is classified correctly by the final majority vote, does not actually lead to greater overfitting. The reason is that using the same type of analysis used to prove Theorem 6.20, one can show that as t_0 increases, not only will the majority vote be correct on each $\mathbf{x} \in S$, but in fact each example will be correctly classified by a $\frac{1}{2} + \gamma'$ fraction of the classifiers, where $\gamma' \to \gamma$ as $t_0 \to \infty$. I.e., the vote is approaching the minimax optimal strategy for the column player in the minimax view given above. This in turn implies that h can be well-approximated over S by a vote of a random sample of $O(1/\gamma^2)$ of its component weak hypotheses h_j . Since these small random majority votes are not overfitting by much, our generalization theorems imply that h cannot be overfitting by much either.

6.11 Stochastic Gradient Descent

We now describe a widely-used algorithm in machine learning, called *stochastic gradient descent* (SGD). The Perceptron algorithm we examined in Section 6.5.3 can be viewed as a special case of this algorithm, as can methods for deep learning.

Let \mathcal{F} be a class of real-valued functions $f_{\mathbf{w}} : \mathbb{R}^d \to \mathbb{R}$ where $\mathbf{w} = (w_1, w_2, \dots, w_n)$ is a vector of parameters. For example, we could think of the class of linear functions where n = d and $f_{\mathbf{w}}(\mathbf{x}) = \mathbf{w}^T x$, or we could have more complicated functions where n > d. For each such function $f_{\mathbf{w}}$ we can define an associated set $h_{\mathbf{w}} = \{\mathbf{x} : f_{\mathbf{w}}(\mathbf{x}) \ge 0\}$, and let $\mathcal{H}_{\mathcal{F}} = \{h_{\mathbf{w}} : f_{\mathbf{w}} \in \mathcal{F}\}$. For example, if \mathcal{F} is the class of linear functions then $\mathcal{H}_{\mathcal{F}}$ is the class of linear separators.

To apply stochastic gradient descent, we also need a loss function $L(f_{\mathbf{w}}(\mathbf{x}), c^*(\mathbf{x}))$ that describes the real-valued penalty we will associate with function $f_{\mathbf{w}}$ for its prediction on an example \mathbf{x} whose true label is $c^*(\mathbf{x})$. The algorithm is then the following:

Stochastic Gradient Descent:

Given: starting point $\mathbf{w} = \mathbf{w}_{init}$ and learning rates $\lambda_1, \lambda_2, \lambda_3, \dots$

(e.g.,
$$\mathbf{w}_{init} = \mathbf{0}$$
 and $\lambda_t = 1$ for all t , or $\lambda_t = 1/\sqrt{t}$)

Consider a sequence of random examples $(\mathbf{x}_1, c^*(\mathbf{x}_1)), (\mathbf{x}_2, c^*(\mathbf{x}_2)), \ldots$

- 1. Given example $(\mathbf{x}_t, c^*(\mathbf{x}_t))$, compute the gradient $\nabla L(f_{\mathbf{w}}(\mathbf{x}_t), c^*(\mathbf{x}_t))$ of the loss of $f_{\mathbf{w}}(\mathbf{x}_t)$ with respect to the weights \mathbf{w} . This is a vector in \mathbb{R}^n whose *i*th component is $\frac{\partial L(f_{\mathbf{w}}(\mathbf{x}_t), c^*(\mathbf{x}_t))}{\partial w_i}$.
- 2. Update: $\mathbf{w} \leftarrow \mathbf{w} \lambda_t \nabla L(f_{\mathbf{w}}(\mathbf{x}_t), c^*(\mathbf{x}_t)).$

Let's now try to understand the algorithm better by seeing a few examples of instantiating the class of functions \mathcal{F} and loss function L.

First, consider n = d and $f_{\mathbf{w}}(\mathbf{x}) = \mathbf{w}^T \mathbf{x}$, so \mathcal{F} is the class of linear predictors. Consider the loss function $L(f_{\mathbf{w}}(\mathbf{x}), c^*(\mathbf{x})) = \max(0, -c^*(\mathbf{x})f_{\mathbf{w}}(\mathbf{x}))$, and recall that $c^*(\mathbf{x}) \in \{-1, 1\}$. In other words, if $f_{\mathbf{w}}(\mathbf{x})$ has the correct sign, then we have a loss of 0, otherwise we have a loss equal to the magnitude of $f_{\mathbf{w}}(\mathbf{x})$. In this case, if $f_{\mathbf{w}}(\mathbf{x})$ has the correct sign and is non-zero, then the gradient will be zero since an infinitesimal change in any of the weights will not change the sign. So, when $h_{\mathbf{w}}(\mathbf{x})$ is correct, the algorithm will leave \mathbf{w} alone. On the other hand, if $f_{\mathbf{w}}(\mathbf{x})$ has the wrong sign, then $\frac{\partial L}{\partial w_i} = -c^*(\mathbf{x})\frac{\partial \mathbf{w} \cdot \mathbf{x}}{\partial w_i} = -c^*(\mathbf{x})x_i$. So, using $\lambda_t = 1$, the algorithm will update $w \leftarrow w + c^*(\mathbf{x})\mathbf{x}$. Note that this is exactly the Perceptron algorithm. (Technically we must address the case that $f_{\mathbf{w}}(\mathbf{x}) = 0$; in this case, we should view $f_{\mathbf{w}}$ as having the wrong sign just barely.)

As a small modification to the above example, consider the same class of linear predictors \mathcal{F} but now modify the loss function to the hinge-loss $L(f_{\mathbf{w}}(\mathbf{x}), c^*(\mathbf{x})) = \max(0, 1 - c^*(\mathbf{x})f_{\mathbf{w}}(\mathbf{x}))$. This loss function now requires $f_{\mathbf{w}}(\mathbf{x})$ to have the correct sign and have magnitude at least 1 in order to be zero. Hinge loss has the useful property that it is an upper bound on error rate: for any sample S, the training error is at most $\sum_{\mathbf{x}\in S} L(f_{\mathbf{w}}(\mathbf{x}), c^*(\mathbf{x}))$. With this loss function, stochastic gradient descent is called the margin perceptron algorithm.

More generally, we could have a much more complex class \mathcal{F} . For example, consider a layered circuit of soft threshold gates. Each node in the circuit computes a linear function of its inputs and then passes this value through an "activation function" such as $a(z) = \tanh(z) = (e^z - e^{-z})/(e^z + e^{-z})$. This circuit could have multiple layers with the output of layer *i* being used as the input to layer i + 1. The vector **w** would be the concatenation of all the weight vectors in the network. This is the idea of *deep neural networks* discussed further in Section 6.13.

While it is difficult to give general guarantees on when stochastic gradient descent will succeed in finding a hypothesis of low error on its training set S, Theorems 6.5 and 6.3

imply that if it does and if S is sufficiently large, we can be confident that its true error will be low as well. Suppose that stochastic gradient descent is run on a machine where each weight is a 64-bit floating point number. This means that its hypotheses can each be described using 64n bits. If S has size at least $\frac{1}{\epsilon}[64n\ln(2) + \ln(1/\delta)]$, by Theorem 6.5 it is unlikely any such hypothesis of true error greater than ϵ will be consistent with the sample, and so if it finds a hypothesis consistent with S, we can be confident its true error is at most ϵ . Or, by Theorem 6.3, if $|S| \ge \frac{1}{2\epsilon^2} (64n\ln(2) + \ln(2/\delta))$ then almost surely the final hypothesis h produced by stochastic gradient descent satisfies true error leas than or equal to training error plus ϵ .

6.12 Combining (Sleeping) Expert Advice

Imagine you have access to a large collection of rules-of-thumb that specify what to predict in different situations. For example, in classifying news articles, you might have one that says "if the article has the word 'football', then classify it as sports" and another that says "if the article contains a dollar figure, then classify it as business". In predicting the stock market, these could be different economic indicators. These predictors might at times contradict each other, e.g., a news article that has both the word "football" and a dollar figure, or a day in which two economic indicators are pointing in different directions. It also may be that no predictor is perfectly accurate with some much better than others. We present here an algorithm for combining a large number of such predictors with the guarantee that if any of them are good, the algorithm will perform nearly as well as each good predictor on the examples on which that predictor fires.

Formally, define a "sleeping expert" to be a predictor h that on any given example **x** either makes a prediction on its label or chooses to stay silent (asleep). We will think of them as black boxes. Now, suppose we have access to n such sleeping experts h_1, \ldots, h_n , and let S_i denote the subset of examples on which h_i makes a prediction (e.g., this could be articles with the word "football" in them). We consider the online learning model, and let mistakes(A, S) denote the number of mistakes of an algorithm A on a sequence of examples S. Then the guarantee of our algorithm A will be that for all i

$$E(mistakes(A, S_i)) \le (1 + \epsilon) \cdot mistakes(h_i, S_i) + O\left(\frac{\log n}{\epsilon}\right)$$

where ϵ is a parameter of the algorithm and the expectation is over internal randomness in the randomized algorithm A.

As a special case, if h_1, \ldots, h_n are concepts from a concept class \mathcal{H} , and so they all make predictions on every example, then A performs nearly as well as the best concept in \mathcal{H} . This can be viewed as a noise-tolerant version of the Halving Algorithm of Section 6.5.2 for the case that no concept in \mathcal{H} is perfect. The case of predictors that make predictions on every example is called the problem of *combining expert advice*, and the more general case of predictors that sometimes fire and sometimes are silent is called the sleeping experts problem.

Combining Sleeping Experts Algorithm:

- Initialize each expert h_i with a weight $w_i = 1$. Let $\epsilon \in (0, 1)$. For each example x, do the following:
- 1. [Make prediction] Let H_x denote the set of experts h_i that make a prediction on x, and let $w_x = \sum_{h_j \in H_x} w_j$. Choose $h_i \in H_x$ with probability $p_{ix} = w_i/w_x$ and predict $h_i(x)$.
- 2. [Receive feedback] Given the correct label, for each $h_i \in H_x$ let $m_{ix} = 1$ if $h_i(x)$ was incorrect, else let $m_{ix} = 0$.
- 3. [Update weights] For each $h_i \in H_x$, update its weight as follows:
 - Let $r_{ix} = \left(\sum_{h_j \in H_x} p_{jx} m_{jx}\right) / (1 + \epsilon) m_{ix}.$
 - Update $w_i \leftarrow w_i (1+\epsilon)^{r_{ix}}$.

Note that $\sum_{h_j \in H_x} p_{jx} m_{jx}$ represents the algorithm's probability of making a mistake on example x. So, h_i is rewarded for predicting correctly $(m_{ix} = 0)$ especially when the algorithm had a high probability of making a mistake, and h_i is penalized for predicting incorrectly $(m_{ix} = 1)$ especially when the algorithm had a low probability of making a mistake.

For each $h_i \notin H_x$, leave w_i alone.

Theorem 6.21 For any set of n sleeping experts h_1, \ldots, h_n , and for any sequence of examples S, the Combining Sleeping Experts Algorithm A satisfies for all i:

$$E(mistakes(A, S_i)) \le (1 + \epsilon) \cdot mistakes(h_i, S_i) + O\left(\frac{\log n}{\epsilon}\right)$$

where $S_i = \{x \in S : h_i \in H_x\}.$

Proof: Consider sleeping expert h_i . The weight of h_i after the sequence of examples S is exactly:

$$w_i = (1+\epsilon)^{\sum_{x \in S_i} \left[\left(\sum_{h_j \in H_x} p_{jx} m_{jx} \right) / (1+\epsilon) - m_{ix} \right]}$$

= $(1+\epsilon)^{E[mistakes(A,S_i)] / (1+\epsilon) - mistakes(h_i,S_i)}.$

Let $w = \sum_{j} w_{j}$. Clearly $w_{i} \leq w$. Therefore, taking logs, we have:

$$E(mistakes(A, S_i))/(1 + \epsilon) - mistakes(h_i, S_i) \leq \log_{1+\epsilon} w.$$

So, using the fact that $\log_{1+\epsilon} w = O(\frac{\log W}{\epsilon})$,

$$E(mistakes(A, S_i)) \leq (1 + \epsilon) \cdot mistakes(h_i, S_i) + O(\frac{\log w}{\epsilon}).$$

Initially, w = n. To prove the theorem, it is enough to prove that w never increases. To do so, we need to show that for each x, $\sum_{h_i \in H_x} w_i (1 + \epsilon)^{r_{ix}} \leq \sum_{h_i \in H_x} w_i$, or equivalently dividing both sides by $\sum_{h_j \in H_x} w_j$ that $\sum_i p_{ix} (1 + \epsilon)^{r_{ix}} \leq 1$, where for convenience we define $p_{ix} = 0$ for $h_i \notin H_x$.

For this we will use the inequalities that for $\beta, z \in [0, 1], \beta^z \leq 1 - (1 - \beta)z$ and $\beta^{-z} \leq 1 + (1 - \beta)z/\beta$. Specifically, we will use $\beta = (1 + \epsilon)^{-1}$. We now have:

$$\begin{split} \sum_{i} p_{ix} (1+\epsilon)^{r_{ix}} &= \sum_{i} p_{ix} \beta^{m_{ix} - (\sum_{j} p_{jx} m_{jx})\beta} \\ &\leq \sum_{i} p_{ix} \left(1 - (1-\beta) m_{ix} \right) \left(1 + (1-\beta) \left(\sum_{j} p_{jx} m_{jx} \right) \right) \\ &\leq \left(\sum_{i} p_{ix} \right) - (1-\beta) \sum_{i} p_{ix} m_{ix} + (1-\beta) \sum_{i} p_{ix} \sum_{j} p_{jx} m_{jx} \\ &= 1 - (1-\beta) \sum_{i} p_{ix} m_{ix} + (1-\beta) \sum_{j} p_{jx} m_{jx} \\ &= 1, \end{split}$$

where the second-to-last line follows from using $\sum_{i} p_{ix} = 1$ in two places. So w never increases and the bound follows as desired.

6.13 Deep learning

Deep learning, or *deep neural networks*, refers to training many-layered networks of nonlinear computational units. The input to the network is an example $\mathbf{x} \in \mathbb{R}^d$. The first layer of the network transforms the example into a new vector $f_1(\mathbf{x})$. Then the second layer transforms $f_1(\mathbf{x})$ into a new vector $f_2(f_1(\mathbf{x}))$, and so on. Finally, the k^{th} layer outputs the final prediction $f_k(f_{k-1}(\dots(f_1(\mathbf{x}))))$. When the learning is supervised the output is typically a vector of probabilities. The motivation for deep learning is that often we are interested in data, such as images, that are given to us in terms of very low-level features, such as pixel intensity values. Our goal is to achieve some higherlevel understanding of each image, such as what objects are in the image and what are they doing. To do so, it is natural to first convert the given low-level representation into one of higher-level features. That is what the layers of the network aim to do. Deep learning is also motivated by multi-task learning, with the idea that a good higher-level representation of data should be useful for a wide range of tasks. Indeed, a common use of deep learning for multi-task learning is to share initial levels of the network across tasks.

A typical architecture of a deep neural network consists of layers of logic units. In a fully connected layer, the output of each gate in the layer is connected to the input of every gate in the next layer. However, if the input is an image one might like to recognize

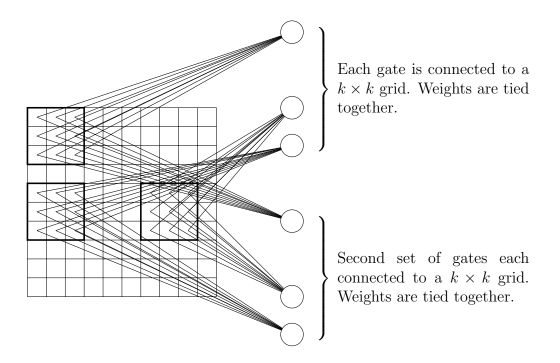


Figure 6.7: Convolution layers

features independent of where they are located in the image. To achieve this one often uses a number of convolution layers. In a convolution layer, each gate gets inputs from a small $k \times k$ grid where k may be 5 to 10. There is a gate for each $k \times k$ square array of the image. The weights on each gate are tied together so that each gate recognizes the same feature. There will be several such collections of gates, so several different features can be learned. Such a level is called a convolution level and the fully connected layers are called autoencoder levels. A technique called *pooling* is used to keep the number of gates reasonable. A small $k \times k$ grid with k typically set to two is used to scan a layer. The stride is set so the grid will provide a non overlapping cover of the layer. Each $k \times k$ input grid will be reduced to a single cell by selecting the maximum input value or the average of the inputs. For k = 2 this reduces the number of cells by a factor of four.

Deep learning networks are trained by stochastic gradient descent (Section 6.11), sometimes called back propagation in the network context. An error function is constructed and the weights are adjusted using the derivative of the error function. This requires that the error function be differentiable. A smooth threshold is used such as

$$\tanh(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}} \quad \text{where} \quad \frac{\partial}{\partial x} \frac{e^e - e^{-e}}{e^x + e^{-x}} = 1 - \left(\frac{e^x - e^{-x}}{e^x + e^{-x}}\right)^2$$

or sigmod $(x) = \frac{1}{1+e^{-x}}$ where

$$\frac{\partial \operatorname{sigmod}(x)}{\partial x} = \frac{e^{-x}}{(1+e^{-x})^2} = \operatorname{sigmod}(x)\frac{e^{-x}}{1+e^{-x}} = \operatorname{sigmoid}(x)\big(1-\operatorname{sigmoid}(x)\big)$$

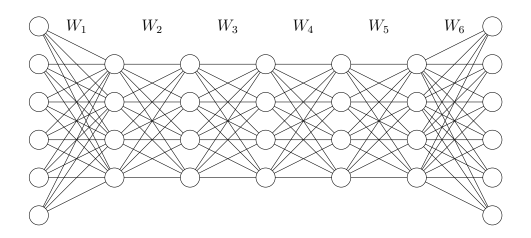


Figure 6.8: A deep learning fully connected network.

In fact the function

$$ReLU(x) = \begin{cases} x & x \ge 0\\ 0 & \text{otherwise} \end{cases} \text{ where } \frac{\partial \text{ReLU}(x)}{\partial x} = \begin{cases} 1 & x \ge 0\\ 0 & \text{otherwise} \end{cases}$$

seems to work well even though typically its derivative at x = 0 is undefined. An advantage of ReLU over sigmoid is that ReLU does not saturate far from the origin.

Training a deep learning network of 7 or 8 levels using gradient descent can be computationally expensive. ²⁶ To address this issue one trains one level at a time on unlabeled data using an idea called autoencoding. There are three levels, the input, a middle level called the hidden level, and an output level as shown in Figure 6.9a. There are two sets of weights. W_1 is the weights of the hidden level gates and W_2 is W_1^T . Let **x** be the input pattern and **y** be the output. The error is $|\mathbf{x} - \mathbf{y}|^2$. One uses gradient descent to reduce the error. Once the weights S_1 are determined they are frozen and a second hidden level of gates is added as in Figure 6.9 b. In this network $W_3 = W_2^T$ and stochastic gradient descent is again used this time to determine W_2 . In this way one level of weights is trained at a time.

The output of the hidden gates is an encoding of the input. An image might be a 10^8 dimensional input and there may only be 10^5 hidden gates. However, the number of images might be 10^7 so even though the dimension of the hidden layer is smaller than the dimension of the input, the number of possible codes far exceeds the number of inputs and thus the hidden layer is a compressed representation of the input. If the hidden layer were the same dimension as the input layer one might get the identity mapping. This does not happen for gradient descent starting with random weights.

 $^{^{26}}$ In the image recognition community, researchers work with networks of 150 levels. The levels tend to be convolution rather than fully connected.

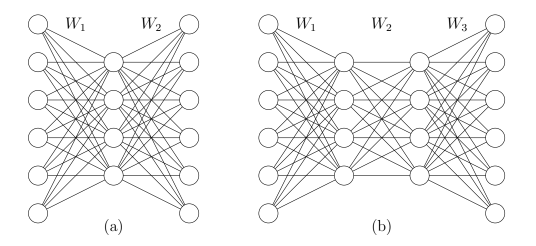


Figure 6.9: Autoencoder technique used to train one level at a time. In the Figure 6.9 (a) train W_1 and W_2 . Then in Figure 6.9 (b), freeze W_1 and train W_2 and W_3 . In this way one trains one set of weights at a time.

The output layer of a deep network typically uses a softmax procedure. Softmax is a generalization of logistic regression where given a set of vectors $\{\mathbf{x_1}, \mathbf{x_2}, \dots, \mathbf{x_n}\}$ with labels $l_1, l_2, \dots, l_n, l_i \in \{0, 1\}$ and with a weight vector \mathbf{w} we define the probability that the label l given x equals 0 or 1 by

$$\operatorname{Prob}(l=1|\mathbf{x}) = \frac{1}{1+e^{-\mathbf{w}^{T}\mathbf{x}}} = \sigma(\mathbf{w}^{T}\mathbf{x})$$

and

$$\operatorname{Prob}(l=0|\mathbf{x}) = 1 - \operatorname{Prob}(l=1/\mathbf{x})$$

where σ is the sigmoid function.

Define a cost function

$$J(\mathbf{w}) = \sum_{i} \left(l_i \log(\operatorname{Prob}(l=1|\mathbf{x})) + (1-l_i) \log(1-\operatorname{Prob}(l=1|\mathbf{x})) \right)$$

and compute \mathbf{w} to minimize $J(\mathbf{x})$. Then

$$J(\mathbf{w}) = \sum_{i} \left(l_i \log(\sigma(\mathbf{w}^{T} \mathbf{x})) + (1 - l_i) \log(1 - \sigma(\mathbf{w}^{T} \mathbf{x})) \right)$$

Since $\frac{\partial \sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x})}{\partial w_j} = \sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x})(1 - \sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x}))x_j$, it follows that $\frac{\partial \log(\sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x}))}{\partial w_j} = \frac{\sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x})(1 - \sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x}))x_j}{\sigma(\mathbf{w}^{\mathbf{T}}\mathbf{x})}$,

Thus

$$\begin{aligned} \frac{\partial J}{\partial w_j} &= \sum_i \left(l_i \frac{\sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x})(1 - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}))}{\sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x})} x_j - (1 - l_i) \frac{(1 - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}))\sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x})}{1 - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x})} x_j \right) \\ &= \sum_i \left(l_i (1 - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x})) x_j - (1 - l_i)\sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}) x_j \right) \\ &= \sum_i \left((l_i x_j - l_i \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}) x_j - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}) x_j + l_i \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}) x_j \right) \\ &= \sum_i \left(l_i - \sigma(\mathbf{w}^{\mathbf{T}} \mathbf{x}) \right) x_j. \end{aligned}$$

Softmax is a generalization of logistic regression to multiple classes. Thus, the labels l_i take on values $\{1, 2, \ldots, k\}$. For an input **x**, softmax estimates the probability of each label. The hypothesis is of the form

$$h_w(x) = \begin{bmatrix} \operatorname{Prob}(l = 1 | \mathbf{x}, \mathbf{w_1}) \\ \operatorname{Prob}(l = 2 | \mathbf{x}, \mathbf{w_2}) \\ \vdots \\ \operatorname{Prob}(l = k | \mathbf{x}, \mathbf{w_k}) \end{bmatrix} = \frac{1}{\sum_{i=1}^k e^{\mathbf{w_i^T x}}} \begin{bmatrix} e^{\mathbf{w_1^T x}} \\ e^{\mathbf{w_2^T x}} \\ \vdots \\ e^{\mathbf{w_k^T x}} \end{bmatrix}$$

where the matrix formed by the weight vectors is

$$W = \begin{bmatrix} \mathbf{w_1} \\ \mathbf{w_2} \\ \vdots \\ \mathbf{w_k} \end{bmatrix}$$

W is a matrix since for each label l_i , there is a vector $\mathbf{w_i}$ of weights.

Consider a set of n inputs $\{\mathbf{x_1}, \mathbf{x_2}, \dots, \mathbf{x_n}\}$. Define

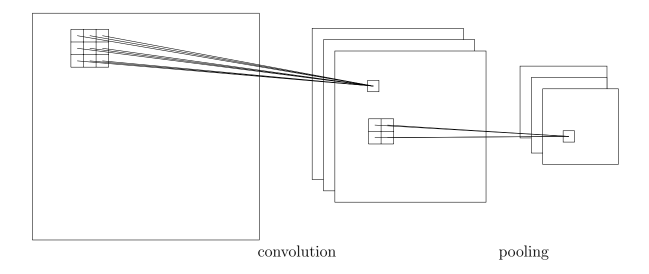
$$\delta(l=k) = \begin{cases} 1 & \text{if } l=k\\ 0 & \text{otherwise} \end{cases}$$

and

$$J(W) = \sum_{i=1}^{n} \sum_{j=1}^{k} \delta(l_i = j) \log \frac{e^{\mathbf{w}_j^{\mathrm{T}} x_i}}{\sum_{h=1}^{k} e^{\mathbf{w}_h^{\mathrm{T}} x_i}}$$

The derivative of the cost function with respect to the weights is

$$\nabla_{\mathbf{w}_{\mathbf{i}}} J(W) = -\sum_{j=1}^{n} \mathbf{x}_{\mathbf{j}} \left(\delta(l_j = k) - \operatorname{Prob}(l_j = k) | \mathbf{x}_{\mathbf{j}}, W \right)$$



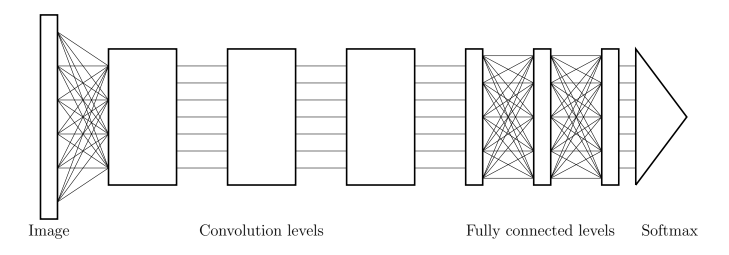


Figure 6.10: A convolution network

Note $\nabla_{\mathbf{w}_{i}} J(W)$ is a vector. Since \mathbf{w}_{i} is a vector, each component of $\nabla_{\mathbf{w}_{i}} J(W)$ is the derivative with respect to one component of the vector \mathbf{w}_{i} .

Over fitting is a major concern in deep learning since large networks can have hundreds of millions of weights. In image recognition, the number of training images can be significantly increased by random jittering of the images. Another technique called *dropout* randomly deletes a fraction of the weights at each training iteration. Regularization is used to assign a cost to the size of weights and many other ideas are being explored.

Deep learning is an active research area. Some of the ideas being explored are what do individual gates or sets of gates learn. If one trains a network twice from starting with random sets of weights, do gates learn the same features? In image recognition, the early convolution layers seem to learn features of images rather than features of the specific set of images they are being trained with. Once a network is trained on say a set of images one of which is a cat one can freeze the weights and then find images that will map to the activation vector generated by the cat image. One can take an artwork image and separate the style from the content and then create an image using the content but a different style []. This is done by taking the activation of the original image and moving it to the manifold of activation vectors of images of a given style. One can do many things of this type. For example one can change the age of a child in an image or change some other feature. []

For more information about deep learning, see $[?]^{.27}$

6.14 Further Current directions

We now briefly discuss a few additional current directions in machine learning, focusing on *semi-supervised* learning, *active* learning, and *multi-task* learning.

6.14.1 Semi-supervised learning

Semi-supervised learning refers to the idea of trying to use a large unlabeled data set U to augment a given labeled data set L in order to produce more accurate rules than would have been achieved using just L alone. The motivation is that in many settings (e.g., document classification, image classification, speech recognition), unlabeled data is much more plentiful than labeled data, so one would like to make use of it if possible. Of course, unlabeled data is missing the labels! Nonetheless it often contains information that an algorithm can take advantage of.

As an example, suppose one believes the target function is a linear separator that separates most of the data by a large margin. By observing enough unlabeled data to estimate the probability mass near to any given linear separator, one could in principle then

²⁷See also the tutorials: http://deeplearning.net/tutorial/deeplearning.pdf and http://deeplearning.stanford.edu/tutorial/.

discard separators in advance that slice through dense regions and instead focus attention on just those that indeed separate most of the distribution by a large margin. This is the high level idea behind a technique known as Semi-Supervised SVMs. Alternatively, suppose data objects can be described by two different "kinds" of features (e.g., a webpage could be described using words on the page itself or using words on links pointing *to* the page), and one believes that each kind should be sufficient to produce an accurate classifier. Then one might want to train a *pair* of classifiers (one on each type of feature) and use unlabeled data for which one is confident but the other is not to bootstrap, labeling such examples with the confident classifier and then feeding them as training data to the less-confident one. This is the high-level idea behind a technique known as Co-Training. Or, if one believes "similar examples should generally have the same label", one might construct a graph with an edge between examples that are sufficiently similar, and aim for a classifier that is correct on the labeled data and has a small cut value on the unlabeled data; this is the high-level idea behind graph-based methods.

A formal model: The batch learning model introduced in Sections 6.1 and 6.3 in essence assumes that one's prior beliefs about the target function be described in terms of a class of functions \mathcal{H} . In order to capture the reasoning used in semi-supervised learning, we need to also describe beliefs about the *relation* between the target function and the data distribution. A clean way to do this is via a *notion of compatibility* χ between a hypothesis h and a distribution \mathcal{D} . Formally, χ maps pairs (h, \mathcal{D}) to [0, 1] with $\chi(h, \mathcal{D}) = 1$ meaning that h is highly compatible with \mathcal{D} and $\chi(h, \mathcal{D}) = 0$ meaning that h is very *in*compatible with \mathcal{D} . The quantity $1 - \chi(h, \mathcal{D})$ is called the *unlabeled error rate* of h, and denoted $err_{unl}(h)$. Note that for χ to be useful, it must be estimatable from a finite sample; to this end, let us further require that χ is an expectation over individual examples. That is, overloading notation for convenience, we require $\chi(h, \mathcal{D}) = \mathbf{E}_{x\sim\mathcal{D}}[\chi(h, x)]$, where $\chi: \mathcal{H} \times \mathcal{X} \to [0, 1]$.

For instance, suppose we believe the target should separate most data by margin γ . We can represent this belief by defining $\chi(h, x) = 0$ if x is within distance γ of the decision boundary of h, and $\chi(h, x) = 1$ otherwise. In this case, $err_{unl}(h)$ will denote the probability mass of \mathcal{D} within distance γ of h's decision boundary. As a different example, in co-training, we assume each example can be described using two "views" that each are sufficient for classification; that is, there exist c_1^*, c_2^* such that for each example $x = \langle x_1, x_2 \rangle$ we have $c_1^*(x_1) = c_2^*(x_2)$. We can represent this belief by defining a hypothesis $h = \langle h_1, h_2 \rangle$ to be compatible with an example $\langle x_1, x_2 \rangle$ if $h_1(x_1) = h_2(x_2)$ and incompatible otherwise; $err_{unl}(h)$ is then the probability mass of examples on which h_1 and h_2 disagree.

As with the class \mathcal{H} , one can either assume that the target is fully compatible (i.e., $err_{unl}(c^*) = 0$) or instead aim to do well as a function of how compatible the target is. The case that we assume $c^* \in \mathcal{H}$ and $err_{unl}(c^*) = 0$ is termed the "doubly realizable case". The concept class \mathcal{H} and compatibility notion χ are both viewed as *known*. **Intuition:** In this framework, the way that unlabeled data helps in learning can be intuitively described as follows. Suppose one is given a concept class \mathcal{H} (such as linear separators) and a compatibility notion χ (such as penalizing h for points within distance γ of the decision boundary). Suppose also that one believes $c^* \in \mathcal{H}$ (or at least is close) and that $err_{unl}(c^*) = 0$ (or at least is small). Then, unlabeled data can help by allowing one to estimate the unlabeled error rate of all $h \in \mathcal{H}$, thereby in principle reducing the search space from \mathcal{H} (all linear separators) down to just the subset of \mathcal{H} that is highly compatible with \mathcal{D} . The key challenge is how this can be done efficiently (in theory, in practice, or both) for natural notions of compatibility, as well as identifying types of compatibility that data in important problems can be expected to satisfy.

A theorem: The following is a semi-supervised analog of our basic sample complexity theorem, Theorem 6.1. First, fix some set of functions \mathcal{H} and compatibility notion χ . Given a labeled sample L, define $\widehat{err}(h)$ to be the fraction of mistakes of h on L. Given an unlabeled sample U, define $\chi(h, U) = \mathbf{E}_{x \sim U}[\chi(h, x)]$ and define $\widehat{err}_{unl}(h) = 1 - \chi(h, U)$. That is, $\widehat{err}(h)$ and $\widehat{err}_{unl}(h)$ are the empirical error rate and unlabeled error rate of h, respectively. Finally, given $\alpha > 0$, define $\mathcal{H}_{\mathcal{D},\chi}(\alpha)$ to be the set of functions $f \in \mathcal{H}$ such that $err_{unl}(f) \leq \alpha$.

Theorem 6.22 If $c^* \in \mathcal{H}$ then with probability at least $1 - \delta$, for labeled set L and unlabeled set U drawn from \mathcal{D} , the $h \in \mathcal{H}$ that optimizes $\widehat{err}_{unl}(h)$ subject to $\widehat{err}(h) = 0$ will have $\operatorname{err}_{\mathcal{D}}(h) \leq \epsilon$ for

$$|U| \ge \frac{2}{\epsilon^2} \left[\ln |\mathcal{H}| + \ln \frac{4}{\delta} \right], \text{ and } |L| \ge \frac{1}{\epsilon} \left[\ln |\mathcal{H}_{\mathcal{D},\chi}(err_{unl}(c^*) + 2\epsilon)| + \ln \frac{2}{\delta} \right].$$

Equivalently, for |U| satisfying this bound, for any |L|, where $h \in \mathcal{H}$ that minimizes $\widehat{err}_{unl}(h)$ subject to $\widehat{err}(h) = 0$ has

$$err_{\mathcal{D}}(h) \leq \frac{1}{|L|} \left[\ln |\mathcal{H}_{\mathcal{D},\chi}(err_{unl}(c^*) + 2\epsilon)| + \ln \frac{2}{\delta} \right]$$

Proof: By Hoeffding bounds, |U| is sufficiently large so that with probability at least $1 - \delta/2$, all $h \in \mathcal{H}$ have $|\widehat{err}_{unl}(h) - err_{unl}(h)| \leq \epsilon$. Thus we have:

$$\{f \in \mathcal{H} : \widehat{err}_{unl}(f) \le err_{unl}(c^*) + \epsilon\} \subseteq \mathcal{H}_{\mathcal{D},\chi}(err_{unl}(c^*) + 2\epsilon).$$

The given bound on |L| is sufficient so that with probability at least $1 - \delta$, all $h \in \mathcal{H}$ with $\widehat{err}(h) = 0$ and $\widehat{err}_{unl}(h) \leq err_{unl}(c^*) + \epsilon$ have $err_{\mathcal{D}}(h) \leq \epsilon$; furthermore, $\widehat{err}_{unl}(c^*) \leq err_{unl}(c^*) + \epsilon$, so such a function h exists. Therefore, with probability at least $1 - \delta$, the $h \in \mathcal{H}$ that optimizes $\widehat{err}_{unl}(h)$ subject to $\widehat{err}(h) = 0$ has $err_{\mathcal{D}}(h) \leq \epsilon$, as desired.

One can view Theorem 6.22 as bounding the number of labeled examples needed to learn well as a function of the "helpfulness" of the distribution \mathcal{D} with respect to χ . Namely, a helpful distribution is one in which $\mathcal{H}_{\mathcal{D},\chi}(\alpha)$ is small for α slightly larger than the compatibility of the true target function, so we do not need much labeled data to identify a good function among those in $\mathcal{H}_{\mathcal{D},\chi}(\alpha)$. For more information on semi-supervised learning, see [?, ?, ?, ?, ?].

6.14.2 Active learning

Active learning refers to algorithms that take an active role in the selection of which examples are labeled. The algorithm is given an initial unlabeled set U of data points drawn from distribution \mathcal{D} and then interactively requests for the labels of a small number of these examples. The aim is to reach a desired error rate ϵ using much fewer labels than would be needed by just labeling random examples (i.e., passive learning).

As a simple example, suppose that data consists of points on the real line and $\mathcal{H} = \{f_a : f_a(x) = 1 \text{ iff } x \geq a\}$ for $a \in R$. That is, \mathcal{H} is the set of all threshold functions on the line. It is not hard to show (see Exercise 6.2) that a random labeled sample of size $O(\frac{1}{\epsilon}\log(\frac{1}{\delta}))$ is sufficient to ensure that with probability $\geq 1 - \delta$, any consistent threshold a' has error at most ϵ . Moreover, it is not hard to show that $\Omega(\frac{1}{\epsilon})$ random examples are necessary for passive learning. However, with active learning we can achieve error ϵ using only $O(\log(\frac{1}{\epsilon}) + \log\log(\frac{1}{\delta}))$ labels. Specifically, first draw an unlabeled sample U of size $O(\frac{1}{\epsilon}\log(\frac{1}{\delta}))$. Then query the leftmost and rightmost points: if these are both negative then output $a' = \infty$, and if these are both positive then output $a' = -\infty$. Otherwise (the leftmost is negative and the rightmost is positive), perform binary search to find two adjacent examples x, x' such that x is negative and x' is positive, and output a' = (x + x')/2. This threshold a' is consistent with the labels on the entire set U, and so by the above argument, has error $\leq \epsilon$ with probability $\geq 1 - \delta$.

The agnostic case, where the target need not belong in the given class \mathcal{H} is quite a bit more subtle, and addressed in a quite general way in the " A^{2} " Agnostic Active learning algorithm [?]. For more information on active learning, see [?, ?].

6.14.3 Multi-task learning

In this chapter we have focused on scenarios where our goal is to learn a single target function c^* . However, there are also scenarios where one would like to learn *multiple* target functions $c_1^*, c_2^*, \ldots, c_n^*$. If these functions are related in some way, then one could hope to do so with less data per function than one would need to learn each function separately. This is the idea of *multi-task learning*.

One natural example is object recognition. Given an image \mathbf{x} , $c_1^*(\mathbf{x})$ might be 1 if \mathbf{x} is a coffee cup and 0 otherwise; $c_2^*(\mathbf{x})$ might be 1 if \mathbf{x} is a pencil and 0 otherwise; $c_3^*(\mathbf{x})$ might be 1 if \mathbf{x} is a laptop and 0 otherwise. These recognition tasks are related in that image features that are good for one task are likely to be helpful for the others as well. Thus, one approach to multi-task learning is to try to learn a common representation under which each of the target functions can be described as a simple function. Another natural example is personalization. Consider a speech recognition system with n different users. In this case there are n target tasks (recognizing the speech of each user) that are clearly related to each other. Some good references for multi-task learning are [?, ?].

6.15 Bibliographic Notes

[TO BE FILLED IN]

6.16 Exercises

Exercise 6.1 (Section 6.2 and 6.3) Consider the instance space $\mathcal{X} = \{0, 1\}^d$ and let \mathcal{H} be the class of 3-CNF formulas. That is, \mathcal{H} is the set of concepts that can be described as a conjunction of clauses where each clause is an OR of up to 3 literals. (These are also called 3-SAT formulas). For example c^* might be $(x_1 \vee \bar{x}_2 \vee x_3)(x_2 \vee x_4)(\bar{x}_1 \vee x_3)(x_2 \vee x_3 \vee x_4)$. Assume we are in the PAC learning setting, so examples are drawn from some underlying distribution D and labeled by some 3-CNF formula c^* .

- 1. Give a number of samples m that would be sufficient to ensure that with probability $\geq 1 \delta$, all 3-CNF formulas consistent with the sample have error at most ϵ with respect to D.
- 2. Give a polynomial-time algorithm for PAC-learning the class of 3-CNF formulas.

Exercise 6.2 (Section 6.2) Consider the instance space $\mathcal{X} = \mathbb{R}$, and the class of functions $\mathcal{H} = \{f_a : f_a(x) = 1 \text{ iff } x \geq a\}$ for $a \in \mathbb{R}$. That is, \mathcal{H} is the set of all threshold functions on the line. Prove that for any distribution \mathcal{D} , a sample S of size $O(\frac{1}{\epsilon}\log(\frac{1}{\delta}))$ is sufficient to ensure that with probability $\geq 1 - \delta$, any $f_{a'}$ such that $\operatorname{err}_S(f_{a'}) = 0$ has $\operatorname{err}_{\mathcal{D}}(f_{a'}) \leq \epsilon$. Note that you can answer this question from first principles, without using the concept of VC-dimension.

Exercise 6.3 (Perceptron; Section 6.5.3) Consider running the Perceptron algorithm in the online model on some sequence of examples S. Let S' be the same set of examples as S but presented in a different order. Does the Perceptron algorithm necessarily make the same number of mistakes on S as it does on S'? If so, why? If not, show such an Sand S' (consisting of the same set of examples in a different order) where the Perceptron algorithm makes a different number of mistakes on S' than it does on S.

Exercise 6.4 (representation and linear separators) Show that any disjunction (see Section 6.3.1) over $\{0,1\}^d$ can be represented as a linear separator. Show that moreover the margin of separation is $\Omega(1/\sqrt{d})$.

Exercise 6.5 (Linear separators; easy) Show that the parity function on $d \ge 2$ Boolean variables cannot be represented by a linear threshold function. The parity function is 1 if and only if an odd number of inputs is 1.

Exercise 6.6 (Perceptron; Section 6.5.3) We know the Perceptron algorithm makes at most $1/\gamma^2$ mistakes on any sequence of examples that is separable by margin γ (we assume all examples are normalized to have length 1). However, it need not find a separator of large margin. If we also want to find a separator of large margin, a natural alternative is to update on any example \mathbf{x} such that $f^*(\mathbf{x})(\mathbf{w} \cdot \mathbf{x}) < 1$; this is called the margin perceptron algorithm.

1. Argue why margin perceptron is equivalent to running stochastic gradient descent on the class of linear predictors $(f_{\mathbf{w}}(\mathbf{x}) = \mathbf{w} \cdot \mathbf{x})$ using hinge loss as the loss function and using $\lambda_t = 1$.

- 2. Prove that on any sequence of examples that are separable by margin γ , this algorithm will make at most $3/\gamma^2$ updates.
- 3. In part 2 you probably proved that each update increases $|\mathbf{w}|^2$ by at most 3. Use this (and your result from part 2) to conclude that if you have a dataset S that is separable by margin γ , and cycle through the data until the margin perceptron algorithm makes no more updates, that it will find a separator of margin at least $\gamma/3$.

Exercise 6.7 (Decision trees, regularization; Section 6.3) Pruning a decision tree: Let S be a labeled sample drawn iid from some distribution \mathcal{D} over $\{0,1\}^n$, and suppose we have used S to create some decision tree T. However, the tree T is large, and we are concerned we might be overfitting. Give a polynomial-time algorithm for pruning T that finds the pruning h of T that optimizes the right-hand-side of Corollary 6.6, i.e., that for a given $\delta > 0$ minimizes:

$$err_S(h) + \sqrt{\frac{\operatorname{size}(h)\ln(4) + \ln(2/\delta)}{2|S|}}$$

To discuss this, we need to define what we mean by a "pruning" of T and what we mean by the "size" of h. A pruning h of T is a tree in which some internal nodes of T have been turned into leaves, labeled "+" or "-" depending on whether the majority of examples in S that reach that node are positive or negative. Let $size(h) = L(h) \log(n)$ where L(h) is the number of leaves in h.

Hint #1: it is sufficient, for each integer L = 1, 2, ..., L(T), to find the pruning of T with L leaves of lowest empirical error on S, that is, $h_L = \operatorname{argmin}_{h:L(h)=L} err_S(h)$. Then you can just plug them all into the displayed formula above and pick the best one.

Hint #2: use dynamic programming.

Exercise 6.8 (Decision trees, sleeping experts; Sections 6.3, 6.12) "Pruning" a Decision Tree Online via Sleeping Experts: Suppose that, as in the above problem, we are given a decision tree T, but now we are faced with a sequence of examples that arrive online. One interesting way we can make predictions is as follows. For each node v of T (internal node or leaf) create two sleeping experts: one that predicts positive on any example that reaches v and one that predicts negative on any example that reaches v. So, the total number of sleeping experts is O(L(T)).

- 1. Say why any pruning h of T, and any assignment of $\{+,-\}$ labels to the leaves of h, corresponds to a subset of sleeping experts with the property that exactly one sleeping expert in the subset makes a prediction on any given example.
- 2. Prove that for any sequence S of examples, and any given number of leaves L, if we run the sleeping-experts algorithm using $\epsilon = \sqrt{\frac{L \log(L(T))}{|S|}}$, then the expected error rate of the algorithm on S (the total number of mistakes of the algorithm divided by

|S|) will be at most $err_S(h_L) + O(\sqrt{\frac{L\log(L(T))}{|S|}})$, where $h_L = \operatorname{argmin}_{h:L(h)=L}err_S(h)$ is the pruning of T with L leaves of lowest error on S.

3. In the above question, we assumed L was given. Explain how we can remove this assumption and achieve a bound of $\min_L \left[err_S(h_L) + O(\sqrt{\frac{L\log(L(T))}{|S|}}) \right]$ by instantiating L(T) copies of the above algorithm (one for each value of L) and then combining these algorithms using the experts algorithm (in this case, none of them will be sleeping).

Exercise 6.9 Kernels; (Section 6.6) Prove Theorem 6.10.

Exercise 6.10 What is the VC-dimension of right corners with axis aligned edges that are oriented with one edge going to the right and the other edge going up?

Exercise 6.11 (VC-dimension; Section 6.9) What is the VC-dimension V of the class \mathcal{H} of axis-parallel boxes in \mathbb{R}^d ? That is, $\mathcal{H} = \{h_{\mathbf{a},\mathbf{b}} : \mathbf{a}, \mathbf{b} \in \mathbb{R}^d\}$ where $h_{\mathbf{a},\mathbf{b}}(\mathbf{x}) = 1$ if $a_i \leq x_i \leq b_i$ for all i = 1, ..., d and $h_{\mathbf{a},\mathbf{b}}(\mathbf{x}) = -1$ otherwise.

- 1. Prove that the VC-dimension is at least your chosen V by giving a set of V points that is shattered by the class (and explaining why it is shattered).
- 2. Prove that the VC-dimension is at most your chosen V by proving that no set of V + 1 points can be shattered.

Exercise 6.12 (VC-dimension, Perceptron, and Margins; Sections 6.5.3, 6.9) Say that a set of points S is shattered by linear separators of margin γ if every labeling of the points in S is achievable by a linear separator of margin at least γ . Prove that no set of $1/\gamma^2 + 1$ points in the unit ball is shattered by linear separators of margin γ .

Hint: think about the Perceptron algorithm and try a proof by contradiction.

Exercise 6.13 (Linear separators) Suppose the instance space \mathcal{X} is $\{0,1\}^d$ and consider the target function c^* that labels an example \mathbf{x} as positive if the least index *i* for which $x_i = 1$ is odd, else labels \mathbf{x} as negative. In other words, $c^*(\mathbf{x}) = \text{``if } x_1 = 1$ then positive else if $x_2 = 1$ then negative else if $x_3 = 1$ then positive else ... else negative.''. Show that the rule can be represented by a linear threshold function.

Exercise 6.14 (Linear separators; harder) Prove that for the problem of Exercise 6.13, we cannot have a linear separator with margin at least 1/f(d) where f(d) is bounded above by a polynomial function of d.

Exercise 6.15 *VC-dimension* Prove that the VC-dimension of circles in the plane is three.

Exercise 6.16 *VC-dimension* Show that the VC-dimension of arbitrary right triangles in the plane is seven.

Exercise 6.17 *VC-dimension* Prove that the VC-dimension of triangles in the plane is seven.

Exercise 6.18 *VC-dimension* Prove that the VC dimension of convex polygons in the plane is infinite.

7 Algorithms for Massive Data Problems: Streaming, Sketching, and Sampling

7.1 Introduction

This chapter deals with massive data problems where the input data is too large to be stored in random access memory. One model for such problems is the streaming model, where n data items a_1, a_2, \ldots, a_n arrive one at a time. For example, the a_i might be IP addresses being observed by a router on the internet. The goal is for our algorithm to compute some statistics, property, or summary of these data items without using too much memory, much less than n. More specifically, we assume each a_i itself is a b-bit quantity where b is not too large; for example, each a_i might be an integer in $\{1, \ldots, m\}$ where $m = 2^b$. Our goal will be to produce some desired output using space polynomial in b and log n; see Figure 7.1.

For example, a very easy problem to solve in the streaming model is to compute the sum of all the a_i . If each a_i is an integer between 1 and $m = 2^b$, then the sum of all the a_i is an integer between 1 and mn and so the number of bits of memory needed to maintain the sum is $O(b + \log n)$. A harder problem, which we discuss shortly, is computing the number of distinct numbers in the input sequence.

One natural approach for tackling a number of problems in the streaming model is to perform random sampling of the input "on the fly". To introduce the basic flavor of sampling on the fly, consider a stream a_1, a_2, \ldots, a_n from which we are to select an index *i* with probability proportional to the value of a_i . When we see an element, we do not know the probability with which to select it since the normalizing constant depends on all of the elements including those we have not yet seen. However, the following method works. Let *S* be the sum of the a_i 's seen so far. Maintain *S* and an index *i* selected with probability $\frac{a_i}{S}$. Initially i = 1 and $S = a_1$. Having seen symbols a_1, a_2, \ldots, a_j , *S* will equal $a_1 + a_2 + \cdots + a_j$ and for each $i \in \{1, \ldots, j\}$, the selected index will be *i* with probability $\frac{a_i}{S}$. On seeing a_{j+1} , change the selected index to j + 1 with probability $\frac{a_{j+1}}{S+a_{j+1}}$ and otherwise keep the same index as before with probability $1 - \frac{a_{j+1}}{S+a_{j+1}}$. If we change the index to j + 1, clearly it was selected with the correct probability. If we keep *i* as our selection, then by induction it will have been selected with probability

$$\left(1 - \frac{a_{j+1}}{S + a_{j+1}}\right)\frac{a_i}{S} = \frac{S}{S + a_{j+1}}\frac{a_i}{S} = \frac{a_i}{S + a_{j+1}}$$

which is the correct probability for selecting index *i*. Finally *s* is updated by adding a_{j+1} to *S*. This problem comes up in many areas such as (sleeping) experts where there is a sequence of weights and we want to pick an expert with probability proportional to its weight. The a_i 's are the weights and the subscript *i* denotes the expert.

stream
$$a_1, a_2, \dots, a_n \longrightarrow$$
 Algorithm some output (low space)

Figure 7.1: High-level representation of the streaming model

7.2 Frequency Moments of Data Streams

An important class of problems concerns the frequency moments of data streams. As mentioned above, a data stream a_1, a_2, \ldots, a_n of length n consists of symbols a_i from an alphabet of m possible symbols which for convenience we denote as $\{1, 2, \ldots, m\}$. Throughout this section, n, m, and a_i will have these meanings and s (for symbol) will denote a generic element of $\{1, 2, \ldots, m\}$. The frequency f_s of the symbol s is the number of occurrences of s in the stream. For a nonnegative integer p, the p^{th} frequency moment of the stream is

$$\sum_{s=1}^{m} f_s^p.$$

Note that the p = 0 frequency moment corresponds to the number of distinct symbols occurring in the stream using the convention $0^0 = 0$. The first frequency moment is just n, the length of the string. The second frequency moment, $\sum_s f_s^2$, is useful in computing the variance of the stream, i.e., the average squared difference from the average frequency.

$$\frac{1}{m}\sum_{s=1}^{m}\left(f_{s}-\frac{n}{m}\right)^{2} = \frac{1}{m}\sum_{s=1}^{m}\left(f_{s}^{2}-2\frac{n}{m}f_{s}+\left(\frac{n}{m}\right)^{2}\right) = \left(\frac{1}{m}\sum_{s=1}^{m}f_{s}^{2}\right) - \frac{n^{2}}{m^{2}}$$

In the limit as p becomes large, $\left(\sum_{s=1}^{m} f_{s}^{p}\right)^{1/p}$ is the frequency of the most frequent element(s).

We will describe sampling based algorithms to compute these quantities for streaming data shortly. But first a note on the motivation for these various problems. The identity and frequency of the the most frequent item, or more generally, items whose frequency exceeds a given fraction of n, is clearly important in many applications. If the items are packets on a network with source and/or destination addresses, the high frequency items identify the heavy bandwidth users. If the data consists of purchase records in a supermarket, the high frequency items are the best-selling items. Determining the number of distinct symbols is the abstract version of determining such things as the number of accounts, web users, or credit card holders. The second moment and variance are useful in networking as well as in database and other applications. Large amounts of network log data are generated by routers that can record the source address, destination address, and the number of packets for all the messages passing through them. This massive data cannot be easily sorted or aggregated into totals for each source/destination. But it is

important to know if some popular source-destination pairs have a lot of traffic for which the variance is one natural measure.

7.2.1 Number of Distinct Elements in a Data Stream

Consider a sequence a_1, a_2, \ldots, a_n of n elements, each a_i an integer in the range 1 to m where n and m are very large. Suppose we wish to determine the number of distinct a_i in the sequence. Each a_i might represent a credit card number extracted from a sequence of credit card transactions and we wish to determine how many distinct credit card accounts there are. Note that this is easy to do in O(m) space by just storing a bit-vector that records which symbols have been seen so far and which have not. It is also easy to do in $O(n \log m)$ space by storing a list of all distinct symbols that have been seen. However, our goal is to use space logarithmic in m and n. We first show that this is impossible using an exact deterministic algorithm. Any deterministic algorithm that determines the number of distinct elements exactly must use at least m bits of memory on some input sequence of length O(m). We then will show how to get around this problem using randomization and approximation.

Lower bound on memory for exact deterministic algorithm

We show that any exact deterministic algorithm must use at least m bits of memory on some sequence of length m + 1. Suppose we have seen a_1, \ldots, a_m , and suppose for sake of contradiction that our algorithm uses less than m bits of memory on all such sequences. There are $2^m - 1$ possible subsets of $\{1, 2, \ldots, m\}$ that the sequence could contain and yet only 2^{m-1} possible states of our algorithm's memory. Therefore there must be two different subsets S_1 and S_2 that lead to the same memory state. If S_1 and S_2 are of different sizes, then clearly this implies an error for one of the input sequences. On the other hand, if they are the same size, then if the next symbol is in S_1 but not S_2 , the algorithm will give the same answer in both cases and therefore must give an incorrect answer on at least one of them.

Algorithm for the Number of distinct elements

Intuition: To beat the above lower bound, look at approximating the number of distinct elements. Our algorithm will produce a number that is within a constant factor of the correct answer using randomization and thus a small probability of failure. First, the idea: suppose the set S of distinct elements was itself chosen uniformly at random from $\{1, \ldots, m\}$. Let *min* denote the minimum element in S. What is the expected value of *min*? If there was one distinct elements, then its expected value would be roughly $\frac{m}{2}$. If there were two distinct elements, the expected value of the minimum would be roughly $\frac{m}{3}$. More generally, for a random set S, the expected value of the minimum is approximately $\frac{m}{|S|+1}$. See Figure 7.2. Solving $min = \frac{m}{|S|+1}$ yields $|S| = \frac{m}{min} - 1$. This suggests keeping track of the minimum element in $O(\log m)$ space and using this equation

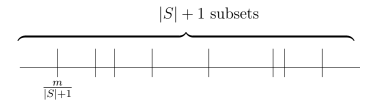


Figure 7.2: Estimating the size of S from the minimum element in S which has value approximately $\frac{m}{|S|+1}$. The elements of S partition the set $\{1, 2, \ldots, m\}$ into |S|+1 subsets each of size approximately $\frac{m}{|S|+1}$.

to give an estimate of |S|.

Converting the intuition into an algorithm via hashing

In general the set S might not have been chosen uniformly at random. If the elements of S were obtained by selecting the |S| smallest elements of $\{1, 2, \ldots, m\}$, the above technique would give a very bad answer. However, we can convert our intuition into an algorithm that works well with high probability on every sequence via hashing. Specifically, we will use a hash function h where

$$h: \{1, 2, \ldots, m\} \to \{0, 1, 2, \ldots, M-1\},\$$

and then instead of keeping track of the minimum element $a_i \in S$, we will keep track of the minimum hash value. The question now is: what properties of a hash function do we need? Since we need to store h, we cannot use a totally random mapping since that would take too many bits. Luckily, a pairwise independent hash function, which can be stored compactly is sufficient.

We recall the formal definition of pairwise independence below. But first recall that a hash function is always chosen at random from a family of hash functions and phrases like "probability of collision" refer to the probability in the choice of hash function.

2-Universal (Pairwise Independent) Hash Functions

A set of hash functions

$$H = \{h \mid h : \{1, 2, \dots, m\} \to \{0, 1, 2, \dots, M - 1\}\}$$

is 2-universal or pairwise independent if for all x and y in $\{1, 2, ..., m\}$ with $x \neq y$, h(x) and h(y) are each equally likely to be any element of $\{0, 1, 2, ..., M-1\}$ and are statistically independent. It follows that a set of hash functions H is 2-universal if and only if for all x and y in $\{1, 2, ..., m\}$, $x \neq y$, h(x) and h(y) are each equally likely to be any element of $\{0, 1, 2, ..., M-1\}$ and for all x and y in $\{1, 2, ..., M-1\}$, and for all w, z we have:

$$\operatorname{Prob}_{h \sim H} \left(h\left(x \right) = w \text{ and } h\left(y \right) = z \right) = \frac{1}{M^2}.$$

We now give an example of a 2-universal family of hash functions. Let M be a prime greater than m. For each pair of integers a and b in the range [0, M - 1], define a hash function

$$h_{ab}\left(x\right) = ax + b \pmod{M}$$

To store the hash function h_{ab} , store the two integers a and b. This requires only $O(\log M)$ space. To see that the family is 2-universal note that h(x) = w and h(y) = z if and only if

$$\begin{pmatrix} x & 1 \\ y & 1 \end{pmatrix} \begin{pmatrix} a \\ b \end{pmatrix} = \begin{pmatrix} w \\ z \end{pmatrix} \pmod{M}$$

If $x \neq y$, the matrix $\begin{pmatrix} x & 1 \\ y & 1 \end{pmatrix}$ is invertible modulo M.²⁸ Thus

$$\binom{a}{b} = \binom{x \ 1}{y \ 1}^{-1} \binom{w}{z} \pmod{M}$$

and for each $\binom{w}{z}$ there is a unique $\binom{a}{b}$. Hence

$$\operatorname{Prob}(h(x) = w \text{ and } h(y) = z) = \frac{1}{M^2}$$

and H is 2-universal.

Analysis of distinct element counting algorithm

Let b_1, b_2, \ldots, b_d be the distinct values that appear in the input. Then the set $S = \{h(b_1), h(b_2), \ldots, h(b_d)\}$ is a set of d random and pairwise independent values from the set $\{0, 1, 2, \ldots, M-1\}$. We now show that $\frac{M}{\min}$ is a good estimate for d, the number of distinct elements in the input, where $\min = \min(S)$.

Lemma 7.1 With probability at least $\frac{2}{3} - \frac{d}{M}$, we have $\frac{d}{6} \leq \frac{M}{\min} \leq 6d$, where min is the smallest element of S.

Proof: First, we show that $\operatorname{Prob}\left(\frac{M}{\min} > 6d\right) < \frac{1}{6} + \frac{d}{M}$. This part does not require pairwise independence.

$$\operatorname{Prob}\left(\frac{M}{\min} > 6d\right) = \operatorname{Prob}\left(\min < \frac{M}{6d}\right) = \operatorname{Prob}\left(\exists k, \ h\left(b_k\right) < \frac{M}{6d}\right)$$
$$\leq \sum_{i=1}^{d} \operatorname{Prob}\left(h(b_i) < \frac{M}{6d}\right) \leq d\left(\frac{\left\lceil \frac{M}{6d} \right\rceil}{M}\right) \leq d\left(\frac{1}{6d} + \frac{1}{M}\right) \leq \frac{1}{6} + \frac{d}{M}.$$

²⁸The primality of M ensures that inverses of elements exist in Z_M^* and M > m ensures that if $x \neq y$, then x and y are not equal mod M.

Next, we show that $\operatorname{Prob}\left(\frac{M}{\min} < \frac{d}{6}\right) < \frac{1}{6}$. This part will use pairwise independence. First, we can write $\operatorname{Prob}\left(\frac{M}{\min} < \frac{d}{6}\right) = \operatorname{Prob}\left(\min > \frac{6M}{d}\right) = \operatorname{Prob}\left(\forall k, h(b_k) > \frac{6M}{d}\right)$. For $i = 1, 2, \ldots, d$ define the indicator variable

$$y_i = \begin{cases} 0 & \text{if } h(b_i) > \frac{6M}{d} \\ 1 & \text{otherwise} \end{cases}$$

and let

$$y = \sum_{i=1}^{d} y_i.$$

We want to show that with good probability, we will see a hash value in $[0, \frac{6M}{d}]$, i.e., that $\operatorname{Prob}(y = 0)$ is small. Now $\operatorname{Prob}(y_i = 1) \geq \frac{6}{d}$, $E(y_i) \geq \frac{6}{d}$, and $E(y) \geq 6$. For 2-way independent random variables, the variance of their sum is the sum of their variances. So $\operatorname{Var}(y) = d\operatorname{Var}(y_1)$. Further, it is easy to see since y_1 is 0 or 1 that $\operatorname{Var}(y_1) = E\left[(y_1 - E(y_1))^2\right] = E(y_1^2) - E^2(y_1) = E(y_1) - E^2(y_1) \leq E(y_1)$. Thus $\operatorname{Var}(y) \leq E(y)$. By the Chebyshev inequality,

$$\operatorname{Prob}\left(\frac{M}{\min} < \frac{d}{6}\right) = \operatorname{Prob}\left(\min > \frac{6M}{d}\right) = \operatorname{Prob}\left(\forall k \ h\left(b_{i}\right) > \frac{6M}{d}\right)$$
$$= \operatorname{Prob}\left(y = 0\right)$$
$$\leq \operatorname{Prob}\left(|y - E\left(y\right)| \ge E\left(y\right)\right)$$
$$\leq \frac{\operatorname{Var}(y)}{E^{2}\left(y\right)} \le \frac{1}{E\left(y\right)} \le \frac{1}{6}$$

Since $\frac{M}{\min} > 6d$ with probability at most $\frac{1}{6} + \frac{d}{M}$ and $\frac{M}{\min} < \frac{d}{6}$ with probability at most $\frac{1}{6}$, $\frac{d}{6} \le \frac{M}{\min} \le 6d$ with probability at least $\frac{2}{3} - \frac{d}{M}$.

7.2.2 Counting the Number of Occurrences of a Given Element.

To count the number of occurrences of a given element in a stream requires at most $\log n$ space where n is the length of the stream. Clearly, for any length stream that occurs in practice, one can afford $\log n$ space. For this reason, the following material may never be used in practice, but the technique is interesting and may give insight into how to solve some other problem.

Consider a string of 0's and 1's of length n in which we wish to count the number of occurrences of 1's. Clearly with $\log n$ bits of memory we could keep track of the exact number of 1's. However, the number can be approximated with only $\log \log n$ bits.

Let *m* be the number of 1's that occur in the sequence. Keep a value *k* such that 2^k is approximately the number of occurrences *m*. Storing *k* requires only $\log \log n$ bits of memory. The algorithm works as follows. Start with k=0. For each occurrence of a 1, add one to *k* with probability $1/2^k$. At the end of the string, the quantity $2^k - 1$ is the

estimate of m. To obtain a coin that comes down heads with probability $1/2^k$, flip a fair coin, one that comes down heads with probability 1/2, k times and report heads if the fair coin comes down heads in all k flips.

Given k, on average it will take 2^k ones before k is incremented. Thus, the expected number of 1's to produce the current value of k is $1 + 2 + 4 + \cdots + 2^{k-1} = 2^k - 1$.

7.2.3 Counting Frequent Elements

The Majority and Frequent Algorithms

First consider the very simple problem of n people voting. There are m candidates, $\{1, 2, \ldots, m\}$. We want to determine if one candidate gets a majority vote and if so who. Formally, we are given a stream of integers a_1, a_2, \ldots, a_n , each a_i belonging to $\{1, 2, \ldots, m\}$, and want to determine whether there is some $s \in \{1, 2, \ldots, m\}$ which occurs more than n/2 times and if so which s. It is easy to see that to solve the problem exactly on read-once streaming data with a deterministic algorithm, requires $\Omega(\min(n, m))$ space. Suppose n is even and the last n/2 items are identical. Suppose also that after reading the first n/2 items, there are two different sets of elements that result in the same content of our memory. In that case, a mistake would occur if the second half of the stream consists solely of an element that is in one set, but not in the other. If $n/2 \ge m$ then there are at least $2^m - 1$ possible subsets of the first n/2 elements. If $n/2 \le m$ then there are $\sum_{i=1}^{n/2} {m \choose i}$ subsets. By the above argument, the number of bits of memory must be at least the base 2 logarithm of the number of subsets, which is $\Omega(\min(m, n))$.

Surprisingly, we can bypass the above lower bound by slightly weakening our goal. Again let's require that if some element appears more than n/2 times, then we must output it. But now, let us say that if no element appears more than n/2 times, then our algorithm may output whatever it wants, rather than requiring that it output "no". That is, there may be "false positives", but no "false negatives".

Majority Algorithm

Store a_1 and initialize a counter to one. For each subsequent a_i , if a_i is the same as the currently stored item, increment the counter by one. If it differs, decrement the counter by one provided the counter is nonzero. If the counter is zero, then store a_i and set the counter to one.

To analyze the algorithm, it is convenient to view the decrement counter step as "eliminating" two items, the new one and the one that caused the last increment in the counter. It is easy to see that if there is a majority element s, it must be stored at the end. If not, each occurrence of s was eliminated; but each such elimination also causes another item to be eliminated. Thus for a majority item not to be stored at the end, more than n items must have eliminated, a contradiction. Next we modify the above algorithm so that not just the majority, but also items with frequency above some threshold are detected. More specifically, the algorithm below finds the frequency (number of occurrences) of each element of $\{1, 2, \ldots, m\}$ to within an additive term of $\frac{n}{k+1}$. That is, for each symbol s, the algorithm produces a value $\tilde{f}_s \in [f_s - \frac{n}{k+1}, f_s]$, where f_s is the true number of occurrences of symbol s in the sequence. It will do so using $O(k \log n + k \log m)$ space by keeping k counters instead of just one counter.

Algorithm Frequent

Maintain a list of items being counted. Initially the list is empty. For each item, if it is the same as some item on the list, increment its counter by one. If it differs from all the items on the list, then if there are less than k items on the list, add the item to the list with its counter set to one. If there are already k items on the list, decrement each of the current counters by one. Delete an element from the list if its count becomes zero.

Theorem 7.2 At the end of Algorithm Frequent, for each $s \in \{1, 2, ..., m\}$, its counter on the list \tilde{f}_s satisfies $\tilde{f}_s \in [f_s - \frac{n}{k+1}, f_s]$. If some s does not occur on the list, its counter is zero and the theorem asserts that $f_s \leq \frac{n}{k+1}$.

Proof: The fact that $\tilde{f}_s \leq f_s$ is immediate. To show $\tilde{f}_s \geq f_s - \frac{n}{k+1}$, view each decrement counter step as eliminating some items. An item is eliminated if the current a_i being read is not on the list and there are already k symbols different from it on the list; in this case, a_i and k other distinct symbols are simultaneously eliminated. Thus, the elimination of each occurrence of an $s \in \{1, 2, \ldots, m\}$ is really the elimination of k + 1 items corresponding to distinct symbols. Thus, no more than n/(k+1) occurrences of any symbol can be eliminated. It is clear that if an item is not eliminated, then it must still be on the list at the end. This proves the theorem.

Theorem 7.2 implies that we can compute the true frequency of every $s \in \{1, 2, ..., m\}$ to within an additive term of $\frac{n}{k+1}$.

7.2.4 The Second Moment

This section focuses on computing the second moment of a stream with symbols from $\{1, 2, \ldots, m\}$. Again, let f_s denote the number of occurrences of symbol s in the stream, and recall that the second moment of the stream is given by $\sum_{s=1}^{m} f_s^2$. To calculate the second moment, for each symbol $s, 1 \leq s \leq m$, independently set a random variable x_s to ± 1 with probability 1/2. In particular, think of x_s as the output of a random hash function h(s) whose range is just the two buckets $\{-1, 1\}$. For now think of h as a fully independent hash function. Maintain a sum by adding x_s to the sum each time the symbol s occurs in the stream. At the end of the stream, the sum will equal $\sum_{s=1}^{m} x_s f_s$. The

expected value of the sum will be zero where the expectation is over the choice of the ± 1 value for the x_s .

$$E\left(\sum_{s=1}^{m} x_s f_s\right) = 0$$

Although the expected value of the sum is zero, its actual value is a random variable and the expected value of the square of the sum is given by

$$E\left(\sum_{s=1}^{m} x_s f_s\right)^2 = E\left(\sum_{s=1}^{m} x_s^2 f_s^2\right) + 2E\left(\sum_{s\neq t} x_s x_t f_s f_t\right) = \sum_{s=1}^{m} f_s^2,$$

The last equality follows since $E(x_s x_t) = E(x_s)E(x_t) = 0$ for $s \neq t$, using pairwise independence of the random variables. Thus

$$a = \left(\sum_{s=1}^{m} x_s f_s\right)^2$$

is an unbiased estimator of $\sum_{s=1}^{m} f_s^2$ in that it has the correct expectation. Note that at this point we could use Markov's inequality to state that $\operatorname{Prob}(a \ge 3\sum_{s=1}^{m} f_s^2) \le 1/3$, but we want to get a tighter guarantee. To do so, consider the second moment of a:

$$E(a^2) = E\left(\sum_{s=1}^m x_s f_s\right)^4 = E\left(\sum_{1 \le s, t, u, v \le m} x_s x_t x_u x_v f_s f_t f_u f_v\right).$$

The last equality is by expansion. Assume that the random variables x_s are 4-wise independent, or equivalently that they are produced by a 4-wise independent hash function. Then, since the x_s are independent in the last sum, if any one of s, u, t, or v is distinct from the others, then the expectation of the whole term is zero. Thus, we need to deal only with terms of the form $x_s^2 x_t^2$ for $t \neq s$ and terms of the form x_s^4 .

Each term in the above sum has four indices, s, t, u, v, and there are $\binom{4}{2}$ ways of choosing two indices that have the same x value. Thus,

$$E(a^{2}) \leq {\binom{4}{2}} E\left(\sum_{s=1}^{m} \sum_{t=s+1}^{m} x_{s}^{2} x_{t}^{2} f_{s}^{2} f_{t}^{2}\right) + E\left(\sum_{s=1}^{m} x_{s}^{4} f_{s}^{4}\right)$$
$$= 6\sum_{s=1}^{m} \sum_{t=s+1}^{m} f_{s}^{2} f_{t}^{2} + \sum_{s=1}^{m} f_{s}^{4}$$
$$\leq 3\left(\sum_{s=1}^{m} f_{s}^{2}\right)^{2} = 3E^{2}(a).$$

Therefore, $Var(a) = E(a^2) - E^2(a) \le 2E^2(a)$.

Since the variance is comparable to the square of the expectation, repeating the process several times and taking the average, gives high accuracy with high probability. Specifically,

Theorem 7.3 If we use $r = \frac{2}{\varepsilon^{2\delta}}$ independently chosen 4-way independent sets of random variables, and let x be the average of the estimates a_1, \ldots, a_r produced, then

$$Prob\left(|x - E(x)| > \varepsilon E(x)\right) < \frac{Var(x)}{\epsilon^2 E(x)} \le \delta.$$

Proof: The proof follows from the fact that taking the average of r independent repetitions reduces variance by a factor of r, so that $Var(x) \leq \delta \varepsilon^2 E(x)$, and then applying Chebyshev's inequality.

It remains to show that we can implement the desired 4-way independent random variables using $O(\log m)$ space. We earlier gave a construction for a pairwise-independent set of hash functions; now we need 4-wise independence, though only into a range of $\{-1, 1\}$. Below we present one such construction.

Error-Correcting codes, polynomial interpolation and limited-way independence

Consider the problem of generating a random *m*-dimensional vector \mathbf{x} of ± 1 's so that any subset of four coordinates is mutually independent. Such an *m*-dimensional vector may be generated from a truly random "seed" of only $O(\log m)$ mutually independent bits. Thus, we need only store the $O(\log m)$ bits and can generate any of the *m* coordinates when needed. For any *k*, there is a finite field *F* with exactly 2^k elements, each of which can be represented with *k* bits and arithmetic operations in the field can be carried out in $O(k^2)$ time. Here, *k* will be the ceiling of $\log_2 m$. A basic fact about polynomial interpolation is that a polynomial of degree at most three is uniquely determined by its value over any field *F* at four points. More precisely, for any four distinct points $a_1, a_2, a_3, a_4 \in F$ and any four possibly not distinct values $b_1, b_2, b_3, b_4 \in F$, there is a unique polynomial $f(x) = f_0 + f_1 x + f_2 x^2 + f_3 x^3$ of degree at most three, so that with computations done over *F*, $f(a_1) = b_1, f(a_2) = b_2, f(a_3) = b_3$, and $f(a_4) = b_4$.

The definition of the pseudo-random ± 1 vector **x** with 4-way independence is simple. Choose four elements f_0, f_1, f_2, f_3 at random from F and form the polynomial $f(s) = f_0 + f_1 s + f_2 s^2 + f_3 s^3$. This polynomial represents **x** as follows. For $s = 1, 2, ..., m, x_s$ is the leading bit of the k-bit representation of f(s). Thus, the *m*-dimensional vector **x** requires only O(k) bits where $k = \lceil \log m \rceil$.

Lemma 7.4 The \mathbf{x} defined above has 4-way independence.

Proof: Assume that the elements of F are represented in binary using ± 1 instead of the traditional 0 and 1. Let s, t, u, and v be any four coordinates of \mathbf{x} and let α, β, γ , and

 δ have values in ± 1 . There are exactly 2^{k-1} elements of F whose leading bit is α and similarly for β , γ , and δ . So, there are exactly $2^{4(k-1)}$ 4-tuples of elements b_1 , b_2 , b_3 , and b_4 in F so that the leading bit of b_1 is α , the leading bit of b_2 is β , the leading bit of b_3 is γ , and the leading bit of b_4 is δ . For each such b_1 , b_2 , b_3 , and b_4 , there is precisely one polynomial f so that $f(s) = b_1$, $f(t) = b_2$, $f(u) = b_3$, and $f(v) = b_4$. The probability that $x_s = \alpha$, $x_t = \beta$, $x_u = \gamma$, and $x_v = \delta$ is precisely

$$\frac{2^{4(k-1)}}{\text{total number of } f} = \frac{2^{4(k-1)}}{2^{4k}} = \frac{1}{16}.$$

Four way independence follows since

$$Prob(x_s = \alpha)Prob(x_t = \beta)Prob(x_u = \gamma)Prob(x_v = \delta)$$
$$= Prob(x_s = \alpha, x_t = \beta, x_u = \gamma \text{ and } x_s = \delta)$$

Lemma 7.4 describes how to get one vector \mathbf{x} with 4-way independence. However, we need $r = O(1/\varepsilon^2)$ vectors. Also the vectors must be mutually independent. Choose r independent polynomials at the outset.

To implement the algorithm with low space, store only the polynomials in memory. This requires $4k = O(\log m)$ bits per polynomial for a total of $O(\frac{\log m}{\varepsilon^2})$ bits. When a symbol s in the stream is read, compute each polynomial at s to obtain the value for the corresponding value of the x_s and update the running sums. x_s is just the leading bit of the polynomial evaluated at s. This calculation requires $O(\log m)$ time. Thus, we repeatedly compute the x_s from the "seeds", namely the coefficients of the polynomials.

This idea of polynomial interpolation is also used in other contexts. Error-correcting codes is an important example. Say we wish to transmit n bits over a channel which may introduce noise. One can introduce redundancy into the transmission so that some channel errors can be corrected. A simple way to do this is to view the n bits to be transmitted as coefficients of a polynomial f(x) of degree n - 1. Now transmit f evaluated at points $1, 2, 3, \ldots, n + m$. At the receiving end, any n correct values will suffice to reconstruct the polynomial and the true message. So up to m errors can be tolerated. But even if the number of errors is at most m, it is not a simple matter to know which values are corrupted. We do not elaborate on this here.

7.3 Matrix Algorithms using Sampling

We now move from the streaming model to a model where the input is stored in memory, but because the input is so large, one would like to produce a much smaller approximation to it, or perform an approximate computation on it in low space. For instance, the input might be stored in a large slow memory and we would like a small "sketch" that can be stored in smaller fast memory and yet retains the important properties of the original input. In fact, one can view a number of results from the chapter on machine learning in this way: we have a large population, and we want to take a small sample, perform some optimization on the sample, and then argue that the optimum solution on the sample will be approximately optimal over the whole population. In the chapter on machine learning, our sample consisted of independent random draws from the overall population or data distribution. Here we will be looking at matrix algorithms and to achieve good multiplicative error bounds rather than additive error bounds, we will perform non-uniform sampling.

Algorithms for matrix problems like matrix multiplication, low-rank approximations, singular value decomposition, compressed representations of matrices, linear regression etc. are widely used. Some of these algorithms take $O(n^3)$ time for $n \times n$ matrices (with improvements to $O(n^{\alpha})$ time for some $\alpha \in (2,3)$, but with worse constants) and so are difficult to carry out for large modern matrices.

The natural alternative to working on the whole input matrix is to pick a random sub-matrix and compute with that. Here, we will pick a subset of columns or rows of the input matrix. If s (for sample size) is the number of columns we are willing to work with, we will do s independent identical trials. In each trial, we select a column of the matrix. All that we have to decide is what the probabilities of picking each column is. A moment's thought will show that sampling uniformly at random is not always good for a good multiplicative error bound. For example, if the input matrix has most columns close to the zero vector and only a few significant columns, then uniformly sampling scolumns, where s is small, will unlikely pick up any of the significant columns; this may have low additive error (if entries are bounded in [-1, 1], say) but the multiplicative error would be huge.

We will see that the "optimal" probabilities are proportional to the squared length of columns. This is referred to as length squared sampling and since its first discovery in the mid-90's, has been proved to have several desirable properties which we will see. Two general notes on this approach:

(i) We will prove error bounds which hold for all input matrices. Our algorithms are randomized (i.e., use a random number generator), so the error bounds are random variables. The bounds are on the expected error (or tail probability bounds on large errors) and apply to any matrix. Note that this contrasts with the situation where we have a stochastic model of the input matrix and only assert error bounds for "most" matrices drawn from the probability distribution of the stochastic model. A mnemoic is - our algorithms can toss coins, but our data does not toss coins. A reason for proving error bounds for any matrix is that in real problems, like the analysis of the web hypertext link matrix or the patient-genome expression matrix, it is the one matrix the user is interested in, not a random matrix. In general, we focus on general algorithms and theorems, not specific applications, so the reader need not be aware of what the two matrices above

mean.

(ii) There is "no free lunch". Since we only work on a small random sample and not on the whole input matrix, our error bounds will not be good for certain matrices. For example, if the input matrix is the identity, it is intuitively clear that picking a few random columns will miss the other directions. Indeed, the initial error bounds we prove using length squared sampling are useful only for "numerically low-rank matrices", which we define later. But there are important applications, for example, Principal Component Analysis, where one has numerically low-rank input matrices and these techniques are useful. There are more sophisticated and time-consuming sampling methods which have error bounds which are good even for non-numerically-low-rank matrices.

To the Reader: Why aren't (i) and (ii) mutually contradictory?

7.3.1 Matrix Multiplication Using Sampling

Suppose A is an $m \times n$ matrix and B is an $n \times p$ matrix and the product AB is desired. We show how to use sampling to get an approximate product faster than the traditional multiplication. Let A(:, k) denote the k^{th} column of A. A(:, k) is a $m \times 1$ matrix. Let B(k, :) be the k^{th} row of B. B(k, :) is a $1 \times n$ matrix. It is easy to see that

$$AB = \sum_{k=1}^{n} A(:,k)B(k,:).$$

Note that for each value of k, A(:, k)B(k, :) is an $m \times p$ matrix each element of which is a single product of elements of A and B. An obvious use of sampling suggests itself. Sample some values for k and compute A(:, k)B(k, :) for the sampled k's and use their suitably scaled sum as the estimate of AB. It turns out that nonuniform sampling probabilities are useful. Define a random variable z that takes on values in $\{1, 2, \ldots, n\}$. Let p_k denote the probability that z assumes the value k. We will solve for a good choice of probabilities later, but for now just consider the p_k as nonnegative numbers that sum to one. Define an associated random matrix variable that has value

$$X = \frac{1}{p_k} A(:,k) B(k,:)$$
(7.1)

with probability p_k . Let E(X) denote the entry-wise expectation.

$$E(X) = \sum_{k=1}^{n} \operatorname{Prob}(z=k) \frac{1}{p_k} A(:,k) B(k,:) = \sum_{k=1}^{n} A(:,k) B(k,:) = AB.$$

This explains the scaling by $\frac{1}{p_k}$ in X. In particular, X is a matrix-valued random variable each of whose components is correct in expectation. We will be interested in

$$E\left(||AB - X||_F^2\right).$$

This can be viewed as the variance of X, defined as the sum of the variances of all its entries.

$$\operatorname{Var}(X) = \sum_{i=1}^{m} \sum_{j=1}^{p} \operatorname{Var}(x_{ij}) = \sum_{ij} E(x_{ij}^{2}) - E(x_{ij})^{2} = \left(\sum_{ij} \sum_{k} p_{k} \frac{1}{p_{k}^{2}} a_{ik}^{2} b_{kj}^{2}\right) - ||AB||_{F}^{2}.$$

We want to choose p_k to minimize this quantity, and notice that we can ignore the $||AB||_F^2$ term since it doesn't depend on the p_k 's at all. We can now simplify by exchanging the order of summations to get

$$\sum_{ij} \sum_{k} p_k \frac{1}{p_k^2} a_{ik}^2 b_{kj}^2 = \sum_{k} \frac{1}{p_k} \left(\sum_{i} a_{ik}^2 \right) \left(\sum_{j} b_{kj}^2 \right) = \sum_{k} \frac{1}{p_k} |A(:,k)|^2 |B(k,:)|^2.$$

What is the best choice of p_k to minimize this sum? It can be seen by calculus²⁹ that the minimizing p_k are proportional to |A(:,k)||B(k,:)|. In the important special case when $B = A^T$, pick columns of A with probabilities proportional to the squared length of the columns. Even in the general case when B is not A^T , doing so simplifies the bounds, so we will use it. This sampling is called "length squared sampling". If p_k is proportional to $|A(:,k)|^2$, i.e., $p_k = \frac{|A(:,k)|^2}{||A||_F^2}$, then

$$E\left(||AB - X||_F^2\right) = \operatorname{Var}(X) \le ||A||_F^2 \sum_k |B(k, :)|^2 = ||A||_F^2 ||B||_F^2.$$

To reduce the variance, we can do s independent trials. Each trial i, i = 1, 2, ..., syields a matrix X_i as in (7.1). We take $\frac{1}{s} \sum_{i=1}^{s} X_i$ as our estimate of AB. Since the variance of a sum of independent random variables is the sum of variances, the variance of $\frac{1}{s} \sum_{i=1}^{s} X_i$ is $\frac{1}{s} \operatorname{Var}(X)$ and so is at most $\frac{1}{s} ||A||_F^2 ||B||_F^2$. Let k_1, \ldots, k_s be the k's chosen in each trial. Expanding this, gives:

$$\frac{1}{s}\sum_{i=1}^{s} X_{i} = \frac{1}{s} \left(\frac{A(:,k_{1}) B(k_{1},:)}{p_{k_{1}}} + \frac{A(:,k_{2}) B(k_{2},:)}{p_{k_{2}}} + \dots + \frac{A(:,k_{s}) B(k_{s},:)}{p_{k_{s}}} \right).$$
(7.2)

We will find it convieneint to write this as the product of an $m \times s$ matrix with a $s \times p$ matrix as follows: Let C be the $m \times s$ matrix consisting of the following columns which are scaled versions of the chosen columns of A:

$$\frac{A(:,k_1)}{\sqrt{sp_{k_1}}}, \frac{A(:,k_2)}{\sqrt{sp_{k_2}}}, \dots \frac{A(:,k_s)}{\sqrt{sp_{k_s}}}$$

Note that the scaling has a nice property (which the reader is asked to verify):

$$E\left(CC^{T}\right) = AA^{T}.\tag{7.3}$$

²⁹By taking derivatives, for any set of nonnegative numbers c_k , $\sum_k \frac{c_k}{p_k}$ is minimized with p_k proportional to $\sqrt{c_k}$.

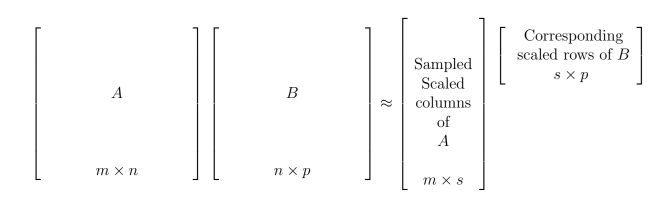


Figure 7.3: Approximate Matrix Multiplication using sampling

Define R to be the $s \times p$ matrix with the corresponding rows of B similarly scaled, namely, R has rows

$$\frac{B(k_1,:)}{\sqrt{sp_{k_1}}}, \frac{B(k_2,:)}{\sqrt{sp_{k_2}}}, \dots \frac{B(k_s,:)}{\sqrt{sp_{k_s}}}.$$

The reader may verify that

$$E\left(R^{T}R\right) = A^{T}A.$$
(7.4)

From (7.2), we see that $\frac{1}{s} \sum_{i=1}^{s} X_i = CR$. This is represented in Figure 7.3. We summarize our discussion in Theorem 7.5.

Theorem 7.5 Suppose A is an $m \times n$ matrix and B is an $n \times p$ matrix. The product AB can be estimated by CR, where C is an $m \times s$ matrix consisting of s columns of A picked according to length-squared distribution and scaled to satisfy (7.3) and R is the $s \times p$ matrix consisting of the corresponding rows of B scaled to satisfy (7.4). The error is bounded by:

$$E\left(||AB - CR||_F^2\right) \le \frac{||A||_F^2 ||B||_F^2}{s}.$$

Thus, to ensure $E(||AB - CR||_F^2) \leq \varepsilon^2 ||A||_F^2 ||B||_F^2$, it suffices to make $s \geq 1/\varepsilon^2$. If $\varepsilon \in \Omega(1)$ (so $s \in O(1)$), then the multiplication CR can be carried out in time O(mp).

When is this error bound good and when is it not? Let's focus on the case that $B = A^T$ so we have just one matrix to consider. If A is the identity matrix, then the guarantee is not very good. In this case, $||AA^T||_F^2 = n$, but the right-hand-side of the inequality is $\frac{n^2}{s}$. So we would need s > n for the bound to be any better than approximating the product with the zero matrix.

More generally, the trivial estimate of zero (all zero matrix) for AA^T makes an error in Frobenius norm of $||AA^T||_F$. What s do we need to ensure that the error is at most this? If $\sigma_1, \sigma_2, \ldots$ are the singular values of A, then the singular values of AA^T are $\sigma_1^2, \sigma_2^2, \ldots$ and

$$||AA^{T}||_{F}^{2} = \sum_{t} \sigma_{t}^{4}$$
 and $||A||_{F}^{2} = \sum_{t} \sigma_{t}^{2}$.

So from the theorem $E(||AA^T - CR||_F^2) \le ||AA^T||_F^2$ provided

$$s \geq \frac{(\sigma_1^2 + \sigma_2^2 + \ldots)^2}{\sigma_1^4 + \sigma_2^4 + \ldots}$$

If rank(A) = r, then there are r non-zero σ_t and the best general upper bound on the ratio $\frac{(\sigma_1^2 + \sigma_2^2 + ...)^2}{\sigma_1^4 + \sigma_2^4 + ...}$ is r, so in general, s needs to be at least r. If A is full rank, this means sampling will not gain us anything over taking the whole matrix!

However, if there is a constant c and a small integer p such that

$$\sigma_1^2 + \sigma_2^2 + \ldots + \sigma_p^2 \ge c(\sigma_1^2 + \sigma_2^2 + \cdots + \sigma_r^2),$$
(7.5)

then,

$$\frac{(\sigma_1^2 + \sigma_2^2 + \ldots)^2}{\sigma_1^4 + \sigma_2^4 + \ldots} \le c^2 \frac{(\sigma_1^2 + \sigma_2^2 + \ldots + \sigma_p^2)^2}{\sigma_1^4 + \sigma_2^4 + \ldots + \sigma_p^2} \le c^2 p,$$

and so $s \ge c^2 p$ gives us a better estimate than the zero matrix. Increasing s by a factor decreases the error by the same factor. The condition (7.5) is indeed the hypothesis of the subject of Principal Component Analysis (PCA) and there are many situations when the data matrix does satisfy the condition and so sampling algorithms are useful.

7.3.2 Implementing Length Squared Sampling in two passes

Traditional matrix algorithms often assume that the input matrix is in Random Access Memory (RAM) and so any particular entry of the matrix can be accessed in unit time. For massive matrices, RAM may be too small to hold the entire matrix, but may be able to hold and compute with the sampled columns and rows.

Consider a high-level model where the input matrix (or matrices) have to be read from "external memory" using a "pass". In one pass, one can read sequentially all entries of the matrix and do some "sampling on the fly".

It is easy to see that two passes suffice to draw a sample of columns of A according to length squared probabilities, even if the matrix is not in row-order or column-order and entries are presented as a linked list. In the first pass, compute the length squared of each column and store this information in RAM. The lengths squared can be computed as running sums. Then, use a random number generator in RAM to determine according to length squared probability the columns to be sampled. Then, make a second pass picking the columns to be sampled.

If the matrix is already presented in external memory in column-order? Then, one pass will do. This is left as an exercise for the reader.

One uses the primitive in Section ??: given a read-once only stream of positive real numbers a_1, a_2, \ldots, a_n , at the end have an $i \in \{1, 2, \ldots, n\}$ with the property that the probability that i was chosen is $\frac{a_i}{\sum_{i=1}^n a_i}$.

7.3.3 Sketch of a Large Matrix

The main result of this section is that for any matrix, a sample of columns and rows, each picked according to length squared distribution provides a good sketch of the matrix. Let A be an $m \times n$ matrix. Pick s columns of A according to length squared distribution. Let C be the $m \times s$ matrix containing the picked columns scaled so as to satisy (7.3), i.e., if A(:,k) is picked, it is scaled by $1/\sqrt{sp_k}$. Similarly, pick r rows of A according to length squared distribution on the rows of A. Let R be the $r \times n$ matrix of the picked rows, scaled as follows: If row k of A is picked, it is scaled by $1/\sqrt{rp_k}$. We then have $E(R^T R) = A^T A$. From C and R, one can find a matrix U so that $A \approx CUR$. The schematic diagram is given in Figure 7.4.

The proof that this is a good approximation makes crucial use of the fact that the sampling of rows and columns is with probability proportional to the squared length. One may recall that the top k singular vectors of the SVD of A, give a similar picture; but the SVD takes more time to compute, requires all of A to be stored in RAM, and does not have the property that the rows and columns are directly from A. The last property, that the approximation involves actual rows/columns of the matrix rather than linear combinations, is called an *interpolative approximation* and is useful in many contexts. However, the SVD yields the best 2-norm approximation. Error bounds for the approximation CUR are weaker.

We briefly touch upon two motivations for such a sketch. Suppose A is the documentterm matrix of a large collection of documents. We are to "read" the collection at the outset and store a sketch so that later, when a query represented by a vector with one entry per term arrives, we can find its similarity to each document in the collection. Similarity is defined by the dot product. In Figure 7.4 it is clear that the matrix-vector product of a query with the right hand side can be done in time O(ns + sr + rm) which would be linear in n and m if s and r are O(1). To bound errors for this process, we need to show that the difference between A and the sketch of A has small 2-norm. Recall that the 2-norm $||A||_2$ of a matrix A is $\max_{|\mathbf{x}|=1} |A\mathbf{x}|$. The fact that the sketch is an interpolative approximation means that our approximation essentially consists a subset of documents and a subset of terms, which may be thought of as a representative set of documents and terms. Additionally, if A is sparse in its rows and columns, each document contains only a small fraction of the terms and each term is in only a small fraction of the documents, then this sparsity property will be preserved in C and R, unlike with SVD.

A second motivation comes from recommendation systems. Here A would be a customer-product matrix whose $(i, j)^{th}$ entry is the preference of customer *i* for prod-

$$A \qquad \left| \approx \begin{bmatrix} Multi \\ plier \end{bmatrix} \begin{bmatrix} Sample rows \\ r \times m \end{bmatrix} \right|$$
$$n \times m \qquad n \times m$$

Figure 7.4: Schematic diagram of the approximation of A by a sample of s columns and r rows.

uct j. The objective is to collect a few sample entries of A and based on them, get an approximation to A so that we can make future recommendations. A few sampled rows of A (preferences of a few customers) and a few sampled columns (customers' preferences for a few products) give a good approximation to A provided that the samples are drawn according to the length-squared distribution.

It remains now to describe how to find U from C and R. There is a $n \times n$ matrix P of the form P = QR that acts as the identity on the space spanned by the rows of R and zeros out all vectors orthogonal to this space. We state this now and postpone the proof.

Lemma 7.6 If RR^T is invertible, then $P = R^T (RR^T)^{-1} R$ has the following properties:

- (i) It acts as the identity matrix on the row space of R. I.e., $P\mathbf{x} = \mathbf{x}$ for every vector \mathbf{x} of the form $\mathbf{x} = R^T \mathbf{y}$ (this defines the row space of R). Furthermore,
- (ii) if \mathbf{x} is orthogonal to the row space of R, then $P\mathbf{x} = \mathbf{0}$.

If RR^T is not invertible, let rank $(RR^T) = r$ and $RR^T = \sum_{t=1}^r \sigma_t \mathbf{u_t} \mathbf{v_t}^T$ be the SVD of RR^T . Then,

$$P = R^T \left(\sum_{t=1}^r \frac{1}{\sigma_t^2} \mathbf{u_t v_t}^T \right) R$$

satisfies (i) and (ii).

We begin with some intuition. In particular, we first present a simpler idea that does not work, but that motivates an idea that does. Write A as AI, where I is the $n \times n$ identity matrix. Approximate the product AI using the algorithm of Theorem 7.5, i.e., by sampling s columns of A according to length-squared. Then, as in the last section, write $AI \approx CW$, where W consists of a scaled version of the s rows of I corresponding to the s columns of A that were picked. Theorem 7.5 bounds the error $||A - CW||_F^2$ by $||A||_F^2||I||_F^2/s = \frac{n}{s}||A||_F^2$. But we would like the error to be a small fraction of $||A||_F^2$ which would require $s \ge n$, which clearly is of no use since this would pick as many or more columns than the whole of A.

Let's use the identity-like matrix P instead of I in the above discussion. Using the fact that R is picked according to length squared sampling, we will show the following proposition later.

Proposition 7.7 $A \approx AP$ and the error $E(||A - AP||_2^2)$ is at most $\frac{1}{\sqrt{r}}||A||_F^2$.

We then use Theorem 7.5 to argue that instead of doing the multiplication AP, we can use the sampled columns of A and the corresponding rows of P. The s sampled columns of A form C. We have to take the corresponding s rows of $P = R^T (RR^T)^{-1}R$, which is the same as taking the corresponding s rows of R^T , and multiplying this by $(RR^T)^{-1}R$. It is easy to check that this leads to an expression of the form CUR. Further, by Theorem 7.5, the error is bounded by

$$E\left(||AP - CUR||_{2}^{2}\right) \le E\left(||AP - CUR||_{F}^{2}\right) \le \frac{||A||_{F}^{2}||P||_{F}^{2}}{s} \le \frac{r}{s}||A||_{F}^{2}, \qquad (7.6)$$

since we will show later that:

Proposition 7.8 $||P||_F^2 \leq r$.

Putting (7.6) and Proposition 7.7 together, and using the fact that by triangle inequality $||A - CUR||_2 \leq ||A - AP||_2 + ||AP - CUR||_2$, which in turn implies that $||A - CUR||_2^2 \leq 2||A - AP||_2^2 + 2||AP - CUR||_2^2$. The main result follows.

Theorem 7.9 Let A be an $m \times n$ matrix and r and s be positive integers. Let C be an $m \times s$ matrix of s columns of A picked according to length squared sampling and let R be a matrix of r rows of A picked according to length squared sampling. Then, we can find from C and R an $s \times r$ matrix U so that

$$E\left(||A - CUR||_{2}^{2}\right) \le ||A||_{F}^{2}\left(\frac{2}{\sqrt{r}} + \frac{2r}{s}\right).$$

If s is fixed, the error is minimized when $r = s^{2/3}$. Choosing $s = r/\varepsilon$ and $r = 1/\varepsilon^2$, the bound becomes $O(\varepsilon)||A||_F^2$. When is this bound meaningful? We discuss this further after first proving all the claims used in the discussion above.

Proof: (of Lemma (7.6)): First for the case when RR^T is invertible. For $\mathbf{x} = R^T \mathbf{y}$, $R^T (RR^T)^{-1}R\mathbf{x} = R^T (RR^T)^{-1}RR^T \mathbf{y} = R^T \mathbf{y} = \mathbf{x}$. If \mathbf{x} is orthogonal to every row of R, then $R\mathbf{x} = \mathbf{0}$, so $P\mathbf{x} = \mathbf{0}$. More generally, if $RR^T = \sum_t \sigma_t \mathbf{u}_t \mathbf{v}_t^T$, then, $R^T \sum_t \frac{1}{\sigma_t^2}R = \sum_t \mathbf{v}_t \mathbf{v}_t^T$ and clearly satisfies (i) and (ii).

Next we prove Proposition 7.7. First, recall that

$$||A - AP||_2^2 = \max_{\{\mathbf{x}:|\mathbf{x}|=1\}} |(A - AP)\mathbf{x}|^2.$$

First suppose \mathbf{x} is in the row space V of R. From Lemma 7.6 $P\mathbf{x} = \mathbf{x}$, so for $\mathbf{x} \in V$, $(A - AP)\mathbf{x} = \mathbf{0}$. Since every vector can be written as a sum of a vector in V plus a vector orthogonal to V, this implies that the maximum must therefore occur at some $\mathbf{x} \in V^{\perp}$. For such \mathbf{x} , by Lemma 7.6, $(A - AP)\mathbf{x} = A\mathbf{x}$. Thus, the question becomes: for unit-length $\mathbf{x} \in V^{\perp}$, how large can $|A\mathbf{x}|^2$ be? To analyze this, write:

$$A\mathbf{x}|^{2} = \mathbf{x}^{T} A^{T} A \mathbf{x} = \mathbf{x}^{T} (A^{T} A - R^{T} R) \mathbf{x} \le ||A^{T} A - R^{T} R||_{2} |\mathbf{x}|^{2} \le ||A^{T} A - R^{T} R||_{2}.$$

This implies that $||A - AP||_2^2 \leq ||A^T A - R^T R||_2$. So, it suffices to prove that $||A^T A - R^T R||_2^2 \leq ||A||_F^4/r$ which follows directly from Theorem 7.5, since we can think of $R^T R$ as a way of estimating $A^T A$ by picking according to length-squared distribution columns of A^T , i.e., rows of A. This proves Proposition 7.7.

Proposition 7.8 is easy to see. By Lemma 7.6, P is the identity on the space V spanned by the rows of R, and $P\mathbf{x} = 0$ for \mathbf{x} perpendicular to the rows of R. Thus $||P||_F^2$ is the sum of its singular values squared which is at most r as claimed.

We now briefly look at the time needed to compute U. The only involved step in computing U is to find $(RR^T)^{-1}$ or do the SVD of RR^T . But note that RR^T is an $r \times r$ matrix and since r is much smaller than n and m, this is fast.

Understanding the bound in Theorem 7.9: To better understand the bound in Theorem 7.9 consider when it is meaningful and when it is not. First, choose parameters $s = \Theta(1/\varepsilon^3)$ and $r = \Theta(1/\varepsilon^2)$ so that the bound becomes $E(||A - CUR||_2^2) \le \varepsilon ||A||_F^2$. Recall that $||A||_F^2 = \sum_i \sigma_i^2(A)$, i.e., the sum of squares of all the singular values of A. Also, for convenience scale A so that $\sigma_1^2(A) = 1$. Then

$$\sigma_1^2(A) = ||A||_2^2 = 1$$
 and $E(||A - CUR||_2^2) \le \varepsilon \sum_i \sigma_i^2(A).$

This, gives an intuitive sense of when the guarantee is good and when it is not. If the top k singular values of A are all $\Omega(1)$ for $k \gg m^{1/3}$, so that $\sum_i \sigma_i^2(A) \gg m^{1/3}$, then the guarantee is only meaningful when $\varepsilon = o(m^{-1/3})$, which is not interesting because it requires s > m. On the other hand, if just the first few singular values of A are large and the rest are quite small, e.g, A represents a collection of points that lie very close to a low-dimensional pancake and in particular if $\sum_i \sigma_i^2(A)$ is a constant, then to be meaningful the bound requires ε to be a small constant. In this case, the guarantee is indeed meaningful because it implies that a constant number of rows and columns provides a good 2-norm approximation to A.

7.4 Sketches of Documents

Suppose one wished to store all the web pages from the WWW. Since there are billions of web pages, one might store just a sketch of each page where a sketch is a few hundred bits that capture sufficient information to do whatever task one had in mind. A web page or a document is a sequence. We begin this section by showing how to sample a set and then how to convert the problem of sampling a sequence into a problem of sampling a set.

Consider subsets of size 1000 of the integers from 1 to 10^6 . Suppose one wished to compute the resemblance of two subsets A and B by the formula

resemblance
$$(A, B) = \frac{|A \cap B|}{|A \cup B|}$$

Suppose that instead of using the sets A and B, one sampled the sets and compared random subsets of size ten. How accurate would the estimate be? One way to sample would be to select ten elements uniformly at random from A and B. However, this method is unlikely to produce overlapping samples. Another way would be to select the ten smallest elements from each of A and B. If the sets A and B overlapped significantly one might expect the sets of ten smallest elements from each of A and B to also overlap. One difficulty that might arise is that the small integers might be used for some special purpose and appear in essentially all sets and thus distort the results. To overcome this potential problem, rename all elements using a random permutation.

Suppose two subsets of size 1000 overlapped by 900 elements. What would the overlap of the 10 smallest elements from each subset be? One would expect the nine smallest elements from the 900 common elements to be in each of the two subsets for an overlap of 90%. The resemblance (A, B) for the size ten sample would be 9/11=0.81.

Another method would be to select the elements equal to zero mod m for some integer m. If one samples mod m the size of the sample becomes a function of n. Sampling mod m allows us to also handle containment.

In another version of the problem one has a sequence rather than a set. Here one converts the sequence into a set by replacing the sequence by the set of all short subsequences of some length k. Corresponding to each sequence is a set of length k subsequences. If k is sufficiently large, then two sequences are highly unlikely to give rise to the same set of subsequences. Thus, we have converted the problem of sampling a sequence to that of sampling a set. Instead of storing all the subsequences, we need only store a small subset of the set of length k subsequences.

Suppose you wish to be able to determine if two web pages are minor modifications of one another or to determine if one is a fragment of the other. Extract the sequence of words occurring on the page. Then define the set of subsequences of k consecutive words from the sequence. Let S(D) be the set of all subsequences of length k occurring in document D. Define resemblance of A and B by

resemblance
$$(A, B) = \frac{|S(A) \cap S(B)|}{|S(A) \cup S(B)|}$$

And define containment as

containment
$$(A, B) = \frac{|S(A) \cap S(B)|}{|S(A)|}$$

Let W be a set of subsequences. Define $\min(W)$ to be the s smallest elements in W and define $\mod(W)$ as the set of elements of w that are zero $\mod m$.

Let π be a random permutation of all length k subsequences. Define F(A) to be the s smallest elements of A and V(A) to be the set mod m in the ordering defined by the permutation.

Then

$$\frac{F(A) \cap F(B)}{F(A) \cup F(B)}$$

and

$$\frac{|V(A) \cap V(B)|}{|V(A) \cup V(B)|}$$

are unbiased estimates of the resemblance of A and B. The value

$$\frac{|V(A) \cap V(B)|}{|V(A)|}$$

is an unbiased estimate of the containment of A in B.

7.5 Bibliography

TO DO

7.6 Exercises

Algorithms for Massive Data Problems

Exercise 7.1 THIS EXERCISE IS IN THE TEXT. SHOULD WE DELETE?

Given a stream of n positive real numbers a_1, a_2, \ldots, a_n , upon seeing

 a_1, a_2, \ldots, a_i keep track of the sum $a = a_1 + a_2 + \cdots + a_i$ and a sample $a_j, j \leq i$ drawn with probability proportional to its value. On reading a_{i+1} , with probability $\frac{a_{i+1}}{a+a_{i+1}}$ replace the current sample with a_{i+1} and update a. Prove that the algorithm selects an a_i from the stream with the probability of picking a_i being proportional to its value.

Exercise 7.2 Given a stream of symbols a_1, a_2, \ldots, a_n , give an algorithm that will select one symbol uniformly at random from the stream. How much memory does your algorithm require?

Exercise 7.3 Give an algorithm to select an a_i from a stream of symbols a_1, a_2, \ldots, a_n with probability proportional to a_i^2 .

Exercise 7.4 How would one pick a random word from a very large book where the probability of picking a word is proportional to the number of occurrences of the word in the book?

Exercise 7.5 Consider a matrix where each element has a probability of being selected. Can you select a row according to the sum of probabilities of elements in that row by just selecting an element according to its probability and selecting the row that the element is in?

Exercise 7.6 For the streaming model give an algorithm to draw s independent samples each with the probability proportional to its value. Justify that your algorithm works correctly.

Frequency Moments of Data Streams Number of Distinct Elements in a Data Stream Lower bound on memory for exact deterministic algorithm Algorithm for the Number of distinct elements Universal Hash Functions

Exercise 7.7 Consider an algorithm that uses a random hash function and gives an estimate of a variable x. Let a be the actual value of x. Suppose that the estimate of x is within $\frac{a}{4} \leq x \leq 4a$ with probability 0.6. The probability of the estimate is with respect to choice of the hash function.

1. How would you improve the estimate of x to $\frac{a}{2} \leq x \leq 2a$ with probability 0.6?

2. How would you improve the probability that $\frac{a}{4} \leq x \leq 4a$ to 0.8?

Exercise 7.8 DELETE? THIS IS IN CURRENT DEFINITION

Show that for a 2-universal hash family $Prob(h(x) = z) = \frac{1}{M+1}$ for all $x \in \{1, 2, ..., m\}$ and $z \in \{0, 1, 2, ..., M\}$.

Exercise 7.9 Let p be a prime. A set of hash functions

 $H = \{h | \{0, 1, \dots, p-1\} \to \{0, 1, \dots, p-1\}\}$

is 3-universal if for all u, v, w, x, y, and z in $\{0, 1, \ldots, p-1\}$, u, v, and w distinct

$$Prob(h(x) = u) = \frac{1}{p}, and$$

 $Prob(h(x) = u, h(y) = v, h(z) = w) = \frac{1}{p^3}$

(a) Is the set $\{h_{ab}(x) = ax + b \mod p \mid 0 \le a, b < p\}$ of hash functions 3-universal?

(b) Give a 3-universal set of hash functions.

Exercise 7.10 Give an example of a set of hash functions that is not 2-universal.

Exercise 7.11 Select a value for k and create a set

$$H = \{ \mathbf{x} | \mathbf{x} = (x_1, x_2, \dots, x_k), x_i \in \{0, 1, \dots, k-1\} \}$$

where the set of vectors H is two way independent and $|H| < k^k$.

Analysis of distinct element counting algorithm Counting the Number of Occurrences of a Given Element.

Exercise 7.12

- (a) What is the variance of the method in Section 7.2.2 of counting the number of occurrences of a 1 with log log n memory?
- (b) Can the algorithm be iterated to use only log log log n memory? What happens to the variance?

Exercise 7.13 Consider a coin that comes down heads with probability p. Prove that the expected number of flips before a head occurs is 1/p.

Exercise 7.14 Randomly generate a string $x_1x_2 \cdots x_n$ of 10^6 0's and 1's with probability $\frac{1}{2}$ of x_i being a 1. Count the number of ones in the string and also estimate the number of ones by the approximate counting algorithm. Repeat the process for p=1/4, 1/8, and 1/16. How close is the approximation?

Counting Frequent Elements The Majority and Frequent Algorithms The Second Moment

Exercise 7.15 Construct an example in which the majority algorithm gives a false positive, *i.e.*, stores a non majority element at the end.

Exercise 7.16 Construct examples where the frequent algorithm in fact does as badly as in the theorem, i.e., it "under counts" some item by n/(k+1).

Exercise 7.17 Recall basic statistics on how an average of independent trials cuts down variance and complete the argument for relative error ε estimate of $\sum_{s=1}^{m} f_s^2$.

Error-Correcting codes, polynomial interpolation and limited-way independence

Exercise 7.18 Let F be a field. Prove that for any four distinct points a_1, a_2, a_3 , and a_4 in F and any four (possibly not distinct) values b_1, b_2, b_3 , and b_4 in F, there is a unique polynomial $f(x) = f_0 + f_1 x + f_2 x^2 + f_3 x^3$ of degree at most three so that $f(a_1) = b_1$, $f(a_2) = b_2$, $f(a_3) = b_3$, and $f(a_4) = b_4$ with all computations done over F.

Sketch of a Large Matrix

Exercise 7.19 Suppose we want to pick a row of a matrix at random where the probability of picking row *i* is proportional to the sum of squares of the entries of that row. How would we do this in the streaming model? Do not assume that the elements of the matrix are given in row order.

- (a) Do the problem when the matrix is given in column order.
- (b) Do the problem when the matrix is represented in sparse notation: it is just presented as a list of triples (i, j, a_{ij}) , in arbitrary order.

Matrix Multiplication Using Sampling

Exercise 7.20 Suppose A and B are two matrices. Prove that $AB = \sum_{k=1}^{n} A(:,k)B(k,:)$.

Exercise 7.21 Generate two 100 by 100 matrices A and B with integer values between 1 and 100. Compute the product AB both directly and by sampling. Plot the difference in L_2 norm between the results as a function of the number of samples. In generating the matrices make sure that they are skewed. One method would be the following. First generate two 100 dimensional vectors a and b with integer values between 1 and 100. Next generate the ith row of A with integer values between 1 and a_i and the ith column of B with integer values between 1 and b_i .

Approximating a Matrix with a Sample of Rows and Columns

Exercise 7.22 Suppose a_1, a_2, \ldots, a_m are nonnegative reals. Show that the minimum of $\sum_{k=1}^{m} \frac{a_k}{x_k}$ subject to the constraints $x_k \ge 0$ and $\sum_k x_k = 1$ is attained when the x_k are proportional to $\sqrt{a_k}$.

Sketches of Documents

Exercise 7.23 Consider random sequences of length n composed of the integers 0 through 9. Represent a sequence by its set of length k-subsequences. What is the resemblance of the sets of length k-subsequences from two random sequences of length n for various values of k as n goes to infinity?

NEED TO CHANGE SUBSEQUENCE TO SUBSTRING

Exercise 7.24 What if the sequences in the Exercise 7.23 were not random? Suppose the sequences were strings of letters and that there was some nonzero probability of a given letter of the alphabet following another. Would the result get better or worse?

Exercise 7.25 Consider a random sequence of length 10,000 over an alphabet of size 100.

- 1. For k = 3 what is probability that two possible successor subsequences for a given subsequence are in the set of subsequences of the sequence?
- 2. For k = 5 what is the probability?

Exercise 7.26 How would you go about detecting plagiarism in term papers?

Exercise 7.27 Suppose you had one billion web pages and you wished to remove duplicates. How would you do this?

Exercise 7.28 Construct two sequences of 0's and 1's having the same set of subsequences of width w.

Exercise 7.29 Consider the following lyrics:

When you walk through the storm hold your head up high and don't be afraid of the dark. At the end of the storm there's a golden sky and the sweet silver song of the lark.

Walk on, through the wind, walk on through the rain though your dreams be tossed and blown. Walk on, walk on, with hope in your heart and you'll never walk alone, you'll never walk alone.

How large must k be to uniquely recover the lyric from the set of all subsequences of symbols of length k? Treat the blank as a symbol.

Exercise 7.30 Blast: Given a long sequence a, say 10^9 and a shorter sequence b, say 10^5 , how do we find a position in a which is the start of a subsequence b' that is close to b? This problem can be solved by dynamic programming but not in reasonable time. Find a time efficient algorithm to solve this problem.

Hint: (Shingling approach) One possible approach would be to fix a small length, say seven, and consider the shingles of a and b of length seven. If a close approximation to b is a substring of a, then a number of shingles of b must be shingles of a. This should allows us to find the approximate location in a of the approximation of b. Some final algorithm should then be able to find the best match.

8 Clustering

8.1 Introduction

Clustering refers to partitioning a set of objects into subsets according to some desired criterion. Often it is an important step in making sense of large amounts of data. Clustering comes up in many contexts. One might want to partition a set of news articles into clusters based on the topics of the articles. Given a set of pictures of people, one might want to group them into clusters based on who is in the image. Or one might want to cluster a set of protein sequences according to the protein function. A related problem is not finding a full partitioning but rather just identifying natural clusters that exist. For example, given a collection of friendship relations among people, one might want to identify any tight-knit groups that exist. In some cases we have a well-defined correct answer, e.g., in clustering photographs of individuals by who is in them, but in other cases the notion of a good clustering may be more subjective.

Before running a clustering algorithm, one first needs to choose an appropriate representation for the data. One common representation is as vectors in \mathbb{R}^d . This corresponds to identifying d real-valued features that are then computed for each data object. For example, to represent documents one might use a "bag of words" representation, where each feature corresponds to a word in the English language and the value of the feature is how many times that word appears in the document. Another common representation is as vertices in a graph, with edges weighted by some measure of how similar or dissimilar the two endpoints are. For example, given a set of protein sequences, one might weight edges based on an edit-distance measure that essentially computes the cost of transforming one sequence into the other. This measure is typically symmetric and satisfies the triangle inequality, and so can be thought of as a finite metric. A point worth noting up front is that often the "correct" clustering of a given set of data depends on your goals. For instance, given a set of photographs of individuals, we might want to cluster the images by who is in them, or we might want to cluster them by facial expression. When representing the images as points in space or as nodes in a weighted graph, it is important that the features we use be relevant to the criterion we care about. In any event, the issue of how best to represent data to highlight the relevant information for a given task is generally addressed using knowledge of the specific domain. From our perspective, the job of the clustering algorithm begins after the data has been represented in some appropriate way.

Not surprisingly, clustering has a long history in terms of algorithmic development. In this chapter, our goals are to (a) discuss some commonly used clustering algorithms and what one can prove about them, and (b) talk more generally about what conditions in terms of what the data looks like are sufficient to be able to produce an approximatelycorrect solution. In the process, we will talk about structural properties of common clustering formulations, as well as relaxations of the clustering goal when there is no clear unique solution. **Preliminaries:** We will follow the standard notation of using n to denote the number of data points and k to denote the number of desired clusters. We will primarily focus on the case that k is known up front, but will also discuss algorithms that produce a sequence of solutions, one for each value of k, as well as algorithms that produce a cluster tree that can encode multiple clusterings at each value of k. We will generally use $A = {\mathbf{a}_1, \ldots, \mathbf{a}_n}$ to denote the n data points.

8.1.1 Two general assumptions on the form of clusters

Before choosing a clustering algorithm, it is useful to have some general idea of what a good clustering should look like. In general, there are two types of assumptions often made that in turn lead to different classes of clustering algorithms.

Center-based clusters: One assumption commonly made is that clusters are *center-based*. This means that the clustering can be defined by k "center points" $\mathbf{c}_1, \ldots, \mathbf{c}_k$, with each data point assigned to whichever center point is closest to it. Note that this assumption does not yet tell us whether one choice of centers is better than another. For this, one needs an objective, or optimization criterion. Three standard criteria often used are k-center, k-median, and k-means clustering, defined as follows.

k-center clustering: Find a partition $C = \{C_1, \ldots, C_k\}$ of A into k clusters, with corresponding centers $\mathbf{c}_1, \ldots, \mathbf{c}_k$, to minimize the maximum distance between any data point and the center of its cluster. That is, we want to minimize

$$\Phi_{kcenter}(\mathcal{C}) = \max_{j=1}^{k} \max_{\mathbf{a}_i \in C_j} d(\mathbf{a}_i, \mathbf{c}_j).$$

k-center clustering makes sense when we believe clusters should be local regions in space. It is also often thought of as the "firehouse location problem" since one can think of it as the problem of locating k fire-stations in a city so as to minimize the maximum distance a fire-truck might need to travel to put out a fire.

k-median clustering: Find a partition $C = \{C_1, \ldots, C_k\}$ of A into k clusters, with corresponding centers $\mathbf{c}_1, \ldots, \mathbf{c}_k$, to minimize the sum of distances between data points and the centers of their clusters. That is, we want to minimize

$$\Phi_{kmedian}(\mathcal{C}) = \sum_{j=1}^{k} \sum_{\mathbf{a}_i \in C_j} d(\mathbf{a}_i, \mathbf{c}_j)$$

k-median clustering is more noise-tolerant than k-center clustering because we are taking a sum rather than a max. A small number of outliers will typically not change the optimal solution by much, unless they are very far away or there are several quite different near-optimal solutions.

k-means clustering: Find a partition $C = \{C_1, \ldots, C_k\}$ of A into k clusters, with corresponding centers $\mathbf{c}_1, \ldots, \mathbf{c}_k$, to minimize the sum of squares of distances between data points and the centers of their clusters. That is, we want to minimize

$$\Phi_{kmeans}(\mathcal{C}) = \sum_{j=1}^{k} \sum_{\mathbf{a}_i \in C_j} d^2(\mathbf{a}_i, \mathbf{c}_j).$$

k-means clustering puts more weight on outliers than k-median clustering, because we are squaring the distances, which magnifies large values. This puts it somewhat in between k-median and k-center clustering in that regard. Using distance squared has some mathematical advantages over using pure distances when data are points in \mathbb{R}^d . For example Corollary 8.2 that asserts that with the distance squared criterion, the optimal center for a given group of data points is its centroid.

The k-means criterion is more often used when data consists of points in \mathbb{R}^d , whereas k-median is more commonly used when we have a finite metric, that is, data are nodes in a graph with distances on edges.

When data are points in \mathbb{R}^d , there are in general two variations of the clustering problem for each of the criteria. We could require that each cluster center be a data point or allow a cluster center to be any point in space. If we require each center to be a data point, the optimal clustering of n data points into k clusters can be solved in time $\binom{n}{k}$ times a polynomial in the length of the data. First, exhaustively enumerate all sets of kdata points as the possible sets of k cluster centers, then associate each point to its nearest center and select the best clustering. No such naive enumeration procedure is available when cluster centers can be any point in space. But, for the k-means problem, Corollary 8.2 shows that once we have identified the data points that belong to a cluster, the best choice of cluster center is the centroid of that cluster, which might not be a data point.

For general values of k, the above optimization problems are all NP-hard.³⁰ So, guarantees on algorithms will typically involve either some form of approximation or some additional assumptions, or both.

High-density clusters: If we do not believe our desired clusters will be center-based, an alternative assumption often made is that clusters consist of high-density regions surrounded by low-density "moats" between them. For example, in the clustering of Figure 8.1 we have one natural cluster A that looks center-based but the other cluster B consists of a ring around cluster A. As seen in the figure, this assumption does not require clusters to correspond to convex regions and it can allow them to be long and stringy. We will examine natural algorithms for clustering data where the desired clusters are believed to be of this form. Note that one difficulty with this assumption is that it can be quite difficult

 $^{^{30}\}mathrm{If}\ k$ is a constant, then as noted above, the version where the centers must be data points can be solved in polynomial time.

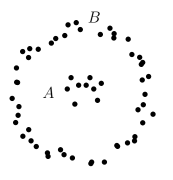


Figure 8.1: Example where the natural clustering is not center-based.

to estimate densities of regions when data lies in high dimensions. So, as a preprocessing step, one may want to first perform some type of projection into a low dimensional space, such as SVD, before running a clustering algorithm.

We begin with a discussion of algorithms for center-based clustering, then examine algorithms for high-density clusters, and then examine some algorithms that allow combining the two. A resource for information on center-based clustering is the book chapter [?].

8.2 k-means Clustering

We assume in this section that data points lie in \mathbb{R}^d and focus on the k-means criterion.

8.2.1 A maximum-likelihood motivation for k-means

We now consider a maximum-likelihood motivation for using the k-means criterion. Suppose that the data was generated according to an equal weight mixture of k spherical well-separated Gaussian densities centered at $\mu_1, \mu_2, \ldots, \mu_k$, each with variance one in every direction. Then the density of the mixture is

Prob(**x**) =
$$\frac{1}{(2\pi)^{d/2}} \frac{1}{k} \sum_{i=1}^{k} e^{-|\mathbf{x}-\boldsymbol{\mu}_i|^2}.$$

Denote by $\boldsymbol{\mu}(\mathbf{x})$ the center nearest to \mathbf{x} . Since the exponential function falls off fast, we can approximate $\sum_{i=1}^{k} e^{-|\mathbf{x}-\boldsymbol{\mu}_i|^2}$ by $e^{-|\mathbf{x}-\boldsymbol{\mu}(\mathbf{x})|^2}$. Thus

$$\operatorname{Prob}(\mathbf{x}) \approx \frac{1}{(2\pi)^{d/2}k} e^{-|\mathbf{x}-\boldsymbol{\mu}(\mathbf{x})|^2}$$

The likelihood of drawing the sample of points $\mathbf{x}_1, \mathbf{x}_2, \ldots, \mathbf{x}_n$ from the mixture, if the centers were $\mu_1, \mu_2, \ldots, \mu_k$, is approximately

$$\frac{1}{k^n} \frac{1}{(2\pi)^{nd/2}} \prod_{i=1}^n e^{-|\mathbf{x}^{(i)} - \boldsymbol{\mu}(\mathbf{x}^{(i)})|^2} = c e^{-\sum_{i=1}^n |\mathbf{x}^{(i)} - \boldsymbol{\mu}(\mathbf{x}^{(i)})|^2}.$$

Minimizing the sum of squared distances to cluster centers finds the maximum likelihood $\mu_1, \mu_2, \ldots, \mu_k$. This motivates using the sum of distance squared to the cluster centers.

8.2.2 Structural properties of the *k*-means objective

Suppose we have already determined the clustering or the partitioning into C_1, C_2, \ldots, C_k . What are the best centers for the clusters? The following lemma shows that the answer is the centroids, the coordinate means, of the clusters.

Lemma 8.1 Let $\{\mathbf{a_1}, \mathbf{a_2}, \ldots, \mathbf{a_n}\}$ be a set of points. The sum of the squared distances of the $\mathbf{a_i}$ to any point \mathbf{x} equals the sum of the squared distances to the centroid of the $\mathbf{a_i}$ plus n times the squared distance from \mathbf{x} to the centroid. That is,

$$\sum_{i} |\mathbf{a}_{i} - \mathbf{x}|^{2} = \sum_{i} |\mathbf{a}_{i} - \mathbf{c}|^{2} + n |\mathbf{c} - \mathbf{x}|^{2}$$

where $\mathbf{c} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{a}_i$ is the centroid of the set of points.

Proof:

$$\sum_{i} |\mathbf{a}_{i} - \mathbf{x}|^{2} = \sum_{i} |\mathbf{a}_{i} - \mathbf{c} + \mathbf{c} - \mathbf{x}|^{2}$$
$$= \sum_{i} |\mathbf{a}_{i} - \mathbf{c}|^{2} + 2(\mathbf{c} - \mathbf{x}) \cdot \sum_{i} (\mathbf{a}_{i} - \mathbf{c}) + n |\mathbf{c} - \mathbf{x}|^{2}$$

Since **c** is the centroid, $\sum_{i} (\mathbf{a}_{i} - \mathbf{c}) = 0$. Thus, $\sum_{i} |\mathbf{a}_{i} - \mathbf{x}|^{2} = \sum_{i} |\mathbf{a}_{i} - \mathbf{c}|^{2} + n |\mathbf{c} - \mathbf{x}|^{2}$

A corollary of Lemma 8.1 is that the centroid minimizes the sum of squared distances since the first term, $\sum_{i} |\mathbf{a}_{i} - \mathbf{c}|^{2}$, is a constant independent of \mathbf{x} and setting $\mathbf{x} = \mathbf{c}$ sets the second term, $n \|\mathbf{c} - \mathbf{x}\|^{2}$, to zero.

Corollary 8.2 Let $\{\mathbf{a_1}, \mathbf{a_2}, \dots, \mathbf{a_n}\}$ be a set of points. The sum of squared distances of the $\mathbf{a_i}$ to a point \mathbf{x} is minimized when \mathbf{x} is the centroid, namely $\mathbf{x} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{a_i}$.

8.2.3 Lloyd's k-means clustering algorithm

Corollary 8.2 suggests the following natural strategy for k-means clustering, known as Lloyd's algorithm. Lloyd's algorithm does not necessarily find a globally optimal solution but will find a locally-optimal one. An important but unspecified step in the algorithm is its initialization: how the starting k centers are chosen. We discuss this after discussing the main algorithm.

Lloyd's algorithm:

Start with k centers.

Cluster each point with the center nearest to it.

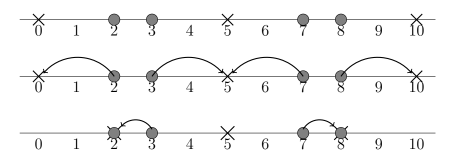
Find the centroid of each cluster and replace the set of old centers with the centroids.

Repeat the above two steps until the centers converge (according to some criterion, such as the k-means score no longer improving).

This algorithm always converges to a local minimum of the objective. To show convergence, we argue that the sum of the squares of the distances of each point to its cluster center always improves. Each iteration consists of two steps. First, consider the step that finds the centroid of each cluster and replaces the old centers with the new centers. By Corollary 8.2, this step improves the sum of internal cluster distances squared. The second step reclusters by assigning each point to its nearest cluster center, which also improves the internal cluster distances.

A problem that arises with some implementations of the k-means clustering algorithm is that one or more of the clusters becomes empty and there is no center from which to measure distance. A simple case where this occurs is illustrated in the following example. You might think how you would modify the code to resolve this issue.

Example: Consider running the k-means clustering algorithm to find three clusters on the following 1-dimension data set: $\{2,3,7,8\}$ starting with centers $\{0,5,10\}$.



The center at 5 ends up with no items and there are only two clusters instead of the desired three.

As noted above, Lloyd's algorithm only finds a local optimum to the k-means objective that might not be globally optimal. Consider, for example, Figure 8.2. Here data lies in three dense clusters in R^2 : one centered at (0, 1), one centered at (0, -1) and one centered at (3, 0). If we initialize with, say, one center at (0, 1) and two centers near (3, 0), then the center at (0, 1) will move to near (0, 0) and capture the points near (0, 1) and (0, -1), whereas the centers near (3, 0) will just stay there, splitting that cluster.

Because the initial centers can substantially influence the quality of the result, there has been significant work on initialization strategies for Lloyd's algorithm. One popular

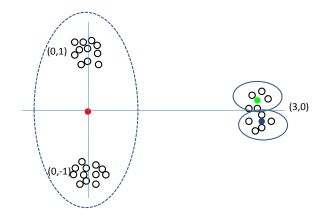


Figure 8.2: A locally-optimal but globally-suboptimal k-means clustering.

strategy is called "farthest traversal". Here, we begin by choosing one data point as initial center \mathbf{c}_1 (say, randomly), then pick the farthest data point from \mathbf{c}_1 to use as \mathbf{c}_2 , then pick the farthest data point from $\{\mathbf{c}_1, \mathbf{c}_2\}$ to use as \mathbf{c}_3 , and so on. These are then used as the initial centers. Notice that this will produce the correct solution in the example in Figure 8.2.

Farthest traversal can unfortunately get fooled by a small number of outliers. To address this, a smoother, probabilistic variation known as k-means++ instead weights data points based on their distance from the previously chosen centers, specifically, proportional to distance squared. Then it selects the next center probabilistically according to these weights. This approach has the nice property that a small number of outliers will not overly influence the algorithm so long as they are not too far away, in which case perhaps they should be their own clusters anyway.

An alternative SVD-based method for initialization is described and analyzed in Section 8.6. Another approach is to run some other approximation algorithm for the k-means problem, and then use its output as the starting point for Lloyd's algorithm. Note that applying Lloyd's algorithm to the output of any other algorithm can only improve its score.

8.2.4 Ward's algorithm

Another popular heuristic for k-means clustering is Ward's algorithm. Ward's algorithm begins with each datapoint in its own cluster, and then repeatedly merges pairs of clusters until only k clusters remain. Specifically, Ward's algorithm merges the two clusters that minimize the immediate increase in k-means cost. That is, for a cluster C, define $cost(C) = \sum_{\mathbf{a}_i \in C} d^2(\mathbf{a}_i, \mathbf{c})$, where **c** is the centroid of C. Then Ward's algorithm merges the pair (C, C') minimizing $cost(C \cup C') - cost(C) - cost(C')$. Thus, Ward's algorithm can be viewed as a greedy k-means algorithm.

8.2.5 k-means clustering on the line

One case where the optimal k-means clustering can be found in polynomial time is when points lie in R^1 , i.e., on the line. This can be done using dynamic programming, as follows.

First, assume without loss of generality that the data points a_1, \ldots, a_n have been sorted, so $a_1 \leq a_2 \leq \ldots \leq a_n$. Now, suppose that for some $i \geq 1$ we have already computed the optimal k'-means clustering for points a_1, \ldots, a_i for all $k' \leq k$; note that this is trivial to do for the base case of i = 1. Our goal is to extend this solution to points a_1, \ldots, a_{i+1} . To do so, we observe that each cluster will contain a consecutive sequence of data points. So, given k', for each $j \leq i+1$, we compute the cost of using a single center for points a_j, \ldots, a_{i+1} , which is the sum of distances of each of these points to their mean value, and then add to that the cost of the optimal k'-1 clustering of points a_1, \ldots, a_{j-1} which we already computed earlier. We store the minimum of these sums, over choices of j, as our optimal k'-means clustering of points a_1, \ldots, a_{i+1} . This has running time of O(kn) for a given value of i, and so overall our running time is $O(kn^2)$.

8.3 k-Center Clustering

In this section, instead of using the k-means clustering criterion, we use the k-center criterion. Recall that the k-center criterion partitions the points into k clusters so as to minimize the maximum distance of any point to its cluster center. Call the maximum distance of any point to its cluster center the radius of the clustering. There is a k-clustering of radius r if and only if there are k spheres, each of radius r, which together cover all the points. Below, we give a simple algorithm to find k spheres covering a set of points. The following lemma shows that this algorithm only needs to use a radius that is at most twice that of the optimal k-center solution. Note that this algorithm is equivalent to the farthest traversal strategy for initializing Lloyd's algorithm.

The Farthest Traversal k-clustering Algorithm

Pick any data point to be the first cluster center. At time t, for t = 2, 3, ..., k, pick the farthest data point from any existing cluster center; make it the t^{th} cluster center.

Theorem 8.3 If there is a k-clustering of radius $\frac{r}{2}$, then the above algorithm finds a k-clustering with radius at most r.

Proof: Suppose for contradiction that there is some data point \mathbf{p} that is distance greater than r from all centers chosen. This means that each new center chosen was distance greater than r from all previous centers, because we could always have chosen \mathbf{p} . This implies that we have k+1 data points, namely the centers chosen plus \mathbf{p} , that are pairwise more than distance r apart. Clearly, no two such points can belong to the same cluster in any k-clustering of radius $\frac{r}{2}$, contradicting the hypothesis.

8.4 Finding Low-Error Clusterings

In the previous sections we saw algorithms for finding a local optimum to the k-means clustering objective, for finding a global optimum to the k-means objective on the line, and for finding a factor 2 approximation to the k-center objective. But what about finding a clustering that is close to the correct answer, such as the true clustering of proteins by function or a correct clustering of news articles by topic? For this we need some assumption about the data and what the correct answer looks like. In the next two sections we will see two different natural assumptions, and algorithms with guarantees based on them.

8.5 Approximation Stability

Implicit in considering objectives like k-means, k-median, or k-center is the hope that the optimal solution to the objective is a desirable clustering. Implicit in considering algorithms that find near-optimal solutions is the hope that near-optimal solutions are also desirable clusterings. Let's now make this idea formal.

Let $C = \{C_1, \ldots, C_k\}$ and $C' = \{C'_1, \ldots, C'_k\}$ be two different k-clusterings of some data set A. A natural notion of the distance between these two clusterings is the fraction of points that would have to be moved between clusters in C to make it match C', where by "match" we allow the indices to be permuted. Since C and C' are both partitions of the set A, this is the same as the fraction of points that would have to be moved among clusters in C' to make it match C. We can write this distance mathematically as:

$$dist(\mathcal{C}, \mathcal{C}') = \min_{\sigma} \frac{1}{n} \sum_{i=1}^{k} |C_i \setminus C'_{\sigma(i)}|,$$

where the minimum is over all permutations σ of $\{1, \ldots, k\}$.

Given an objective Φ (such as k-means, k-median, etc), define \mathcal{C}^* to be the clustering that minimizes Φ . Define \mathcal{C}_T to be the "target" clustering we are aiming for, such as correctly clustering documents by topic or correctly clustering protein sequences by their function. For $c \geq 1$ and $\epsilon > 0$ we say that a data set satisfies (c, ϵ) approximationstability with respect to objective Φ if every clustering \mathcal{C} with $\Phi(\mathcal{C}) \leq c \cdot \Phi(\mathcal{C}^*)$ satisfies $dist(\mathcal{C}, \mathcal{C}_T) < \epsilon$. That is, it is sufficient to be within a factor c of optimal to the objective Φ in order for the fraction of points clustered incorrectly to be less than ϵ .

What is interesting about approximation-stability is the following. The current best polynomial-time approximation guarantee known for the k-means objective is roughly a factor of nine, and for k-median it is roughly a factor 2.7; beating a factor 1 + 3/e for k-means and 1 + 1/e for k-median are both NP-hard. Nonetheless, given data that satisfies $(1.1, \epsilon)$ approximation-stability for the k-median objective, it turns out that so long as ϵn is sufficiently small compared to the smallest cluster in C_T , we can efficiently find a clustering that is ϵ -close to C_T . That is, we can perform as well as if we had a generic 1.1-factor approximation algorithm, even though achieving a 1.1-factor approximation is NP-hard in general. Results known for the k-means objective are somewhat weaker, finding a clustering that is $O(\epsilon)$ -close to C_T . Here, we show this for the k-median objective where the analysis is cleanest. We make a few additional assumptions in order to focus on the main idea. In the following one should think of ϵ as $o(\frac{c-1}{k})$. The results described here apply to data in any metric space, it need not be R^d .³¹

For simplicity and ease of notation assume that $C_T = C^*$; that is, the target clustering is also the optimum for the objective. For a given data point \mathbf{a}_i , define its *weight* $w(\mathbf{a}_i)$ to be its distance to the center of its cluster in C^* . Notice that for the k-median objective, we have $\Phi(C^*) = \sum_{i=1}^n w(\mathbf{a}_i)$. Define $w_{avg} = \Phi(C^*)/n$ to be the average weight of the points in A. Finally, define $w_2(\mathbf{a}_i)$ to be the distance of \mathbf{a}_i to its second-closest center in C^* . We now begin with a useful lemma.

Lemma 8.4 Assume dataset A satisfies (c, ϵ) approximation-stability with respect to the k-median objective, each cluster in C_T has size at least $2\epsilon n$, and $C_T = C^*$. Then,

- 1. Fewer than ϵn points \mathbf{a}_i have $w_2(\mathbf{a}_i) w(\mathbf{a}_i) \leq (c-1)w_{avg}/\epsilon$.
- 2. At most $5\epsilon n/(c-1)$ points \mathbf{a}_i have $w(\mathbf{a}_i) \ge (c-1)w_{avg}/(5\epsilon)$.

Proof: For part (1), suppose that ϵn points \mathbf{a}_i have $w_2(\mathbf{a}_i) - w(\mathbf{a}_i) \leq (c-1)w_{avg}/\epsilon$. Consider modifying \mathcal{C}_T to a new clustering \mathcal{C}' by moving each of these points \mathbf{a}_i into the cluster containing its second-closest center. By assumption, the k-means cost of the clustering has increased by at most $\epsilon n(c-1)w_{avg}/\epsilon = (c-1)\Phi(\mathcal{C}^*)$. This means that $\Phi(\mathcal{C}') \leq c \cdot \Phi(\mathcal{C}^*)$. However, $dist(\mathcal{C}', \mathcal{C}_T) = \epsilon$ because (a) we moved ϵn points to different clusters, and (b) each cluster in \mathcal{C}_T has size at least $2\epsilon n$ so the optimal permutation σ in the definition of dist remains the identity. So, this contradicts approximation stability. Part (2) follows from the definition of "average"; if it did not hold then $\sum_{i=1}^n w(\mathbf{a}_i) > nw_{avg}$, a contradiction.

A datapoint \mathbf{a}_i is *bad* if it satisfies either item (1) or (2) of Lemma 8.4 and *good* if it satisfies neither one. So, there are at most $b = \epsilon n + \frac{5\epsilon n}{c-1}$ bad points and the rest are good. Define "critical distance" $d_{crit} = \frac{(c-1)w_{avg}}{5\epsilon}$. So, Lemma 8.4 implies that the good points have distance at most d_{crit} to the center of their own cluster in \mathcal{C}^* and distance at least $5d_{crit}$ to the center of any other cluster in \mathcal{C}^* .

This suggests the following algorithm. Suppose we create a graph G with the points \mathbf{a}_i as vertices, and edges between any two points \mathbf{a}_i and \mathbf{a}_j with $d(\mathbf{a}_i, \mathbf{a}_j) < 2d_{crit}$. Notice

³¹An example of a data set satisfying (2, 0.2/k) approximation stability would be k clusters of n/k points each, where in each cluster, 90% of the points are within distance 1 of the cluster center (call this the "core" of the cluster), with the other 10% arbitrary, and all cluster cores are at distance at least 10k apart.

that by triangle inequality, the good points within the same cluster in C^* have distance less than $2d_{crit}$ from each other so they will be fully connected and form a clique. Also, again by triangle inequality, any edge that goes between different clusters must be between two bad points. In particular, if \mathbf{a}_i is a good point in one cluster, and it has an edge to some other point \mathbf{a}_j , then \mathbf{a}_j must have distance less than $3d_{crit}$ to the center of \mathbf{a}_i 's cluster. This means that if \mathbf{a}_j had a different closest center, which obviously would also be at distance less than $3d_{crit}$, then \mathbf{a}_i would have distance less than $2d_{crit} + 3d_{crit} = 5d_{crit}$ to that center, violating its goodness. So, bridges in G between different clusters can only occur between bad points.

Assume now that each cluster in C_T has size at least 2b+1; this is the sense in which we are requiring that ϵn be small compared to the smallest cluster in C_T . In this case, create a new graph H by connecting any two points \mathbf{a}_i and \mathbf{a}_j that share at least b+1 neighbors in common in G, themselves included. Since every cluster has at least 2b+1-b=b+1good points, and these points are fully connected in G, this means that H will contain an edge between every pair of good points in the same cluster. On the other hand, since the only edges in G between different clusters are between bad points, and there are at most b bad points, this means that H will not have any edges between different clusters in C_T . Thus, if we take the k largest connected components in H, these will all correspond to subsets of different clusters in C_T , with at most b points remaining.

At this point we have a correct clustering of all but at most b points in A. Call these clusters C_1, \ldots, C_k , where $C_j \subseteq C_j^*$. To cluster the remaining points \mathbf{a}_i , we assign them to the cluster C_j that minimizes the median distance between \mathbf{a}_i and points in C_j . Since each C_j has more good points than bad points, and each good point in C_j has distance at most d_{crit} to center \mathbf{c}_j^* , by triangle inequality the median of these distances must lie in the range $[d(\mathbf{a}_i, \mathbf{c}_i^*) - d_{crit}, d(\mathbf{a}_i, \mathbf{c}_i^*) + d_{crit}]$. This means that this second step will correctly cluster all points \mathbf{a}_i for which $w_2(\mathbf{a}_i) - w(\mathbf{a}_i) > 2d_{crit}$. In particular, we correctly cluster all points except possibly for some of the at most ϵn satisfying item (1) of Lemma 8.4.

The above discussion assumes the value d_{crit} is known to our algorithm; we leave it as an exercise to the reader to modify the algorithm to remove this assumption. Summarizing, we have the following algorithm and theorem.

Algorithm k-Median Stability (given c, ϵ, d_{crit})

- 1. Create a graph G with a vertex for each datapoint in A, and an edge between vertices i and j if $d(\mathbf{a}_i, \mathbf{a}_j) \leq 2d_{crit}$.
- 2. Create a graph H with a vertex for each vertex in G and an edge between vertices i and j if i and j share at least b + 1 neighbors in common, themselves included, for $b = \epsilon n + \frac{5\epsilon n}{c-1}$. Let C_1, \ldots, C_k denote the k largest connected components in H.
- 3. Assign each point not in $C_1 \cup \ldots \cup C_k$ to the cluster C_j of smallest median distance.

Theorem 8.5 Assume A satisfies (c, ϵ) approximation-stability with respect to the kmedian objective, that each cluster in C_T has size at least $\frac{10\epsilon}{c-1}n+2\epsilon n+1$, and that $C_T = C^*$. Then Algorithm k-Median Stability will find a clustering C such that $dist(C, C_T) \leq \epsilon$.

8.6 Spectral Clustering

Spectral clustering is used in many applications and the technique varies depending on the application. If one is clustering the rows of a matrix where the matrix elements are real numbers, then one selects a value for k and computes the top k-singular vectors. The rows of the matrix are projected onto the space spanned by the top k-singular vectors and k-means clustering is used to find k clusters in this lower dimensional space.

If one is clustering the rows of the adjacency matrix A of an undirected graph, a different techniques is used. One uses the Laplacian, L = D - A, where D is a diagonal matrix with degrees of the vertices on the diagonal. Since each row of the Laplacian sums to zero, the all ones vector is a singular vector with singular value zero. The Laplacian is positive semi definite and thus all singular values are non negative. To see this, express L as $L = EE^T$ where E is a matrix whose rows correspond to vertices and whose columns correspond to edges in the graph. Each column of E has two entries corresponding to the two vertices the edge connects. One entry is +1 the other -1. Then for any $\mathbf{x}, \mathbf{x}^T L \mathbf{x} = \mathbf{x}^T E E^T \mathbf{x} = |E\mathbf{x}|^2 \geq 0$. Two normalized versions of the Laplacian are also used. One version is the symmetric normalized Laplacian $L_{sym} = D^{-\frac{1}{2}} L D^{-\frac{1}{2}}$ and another is a normalized version, $L_{rw} = D^{-1}L$ that corresponds to a random walk in the graph. n clustering vertices of a graph one just clusters the rows of the matrix whose columns are the singular vectors of the appropriate version of the Laplacian.

The Laplacian L = D - A is often used to partition a graph into two approximately equal size pieces with a small number of edges between the two pieces. Mathematically this means finding a vector **v** of half ones and half minus ones where a one says a vertex is on one side of the partition and a minus one says it is on the other side. Note that L**v** is a vector whose i^{th} coordinate equals minus two times the number of edges from vertex *i* to vertices on the other side of the partition. It we minimize the sum of the squares of the number of edges from each vertex on one side of the partition. We are almost finding the second singular vector. Requiring *v* to be half ones and half minus ones makes *v* perpendicular to the first singular vector of all ones. If we were not requiring *v* to consist of ones and minus ones we would be calculating

$$v_2 = \arg \max_{v \perp v_1} |Av|^2,$$

the second singular vector. If one calculates the second singular vector of the graph Laplacian and partitions the vertices by the sign of the corresponding coordinate of the second singular vector, they should get a partition of roughly equal size components with a small number of edges between the two blocks of the partition.

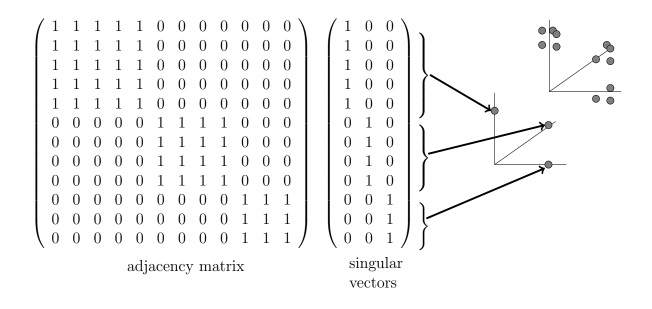


Figure 8.3: In spectral clustering the vertices of a graph, one finds a few top singular vectors of the adjacency matrix and forms a new matrix whose columns are these singular vectors. The rows of this matrix are points in a lower dimensional space. These points can be clustered with k-means cluster. In the above example there are three clusters that are cliques with no connecting vertices. If one removed a few edges from the cliques and added a few edges connecting vertices in different cliques, the rows of a clique would map to clusters of points as shown in the top right rather than all to a single point.

Sometimes there is clearly a correct target clustering we want to find, but approximately optimal k-means or k-median clusterings can be very far from the target. We describe two important stochastic models where this situation arises. In these cases as well as many others, a clustering algorithm based on SVD, called Spectral Clustering is useful.

Spectral Clustering first projects data points onto the space spanned by the top k singular vectors of the data matrix and works in the projection. It is widely used and indeed finds a clustering close to the target clustering for data arising from many stochastic models. We will prove this in a generalized setting which includes both stochastically generated data and data with no stochastic models.

8.6.1 Stochastic Block Model

Stochastic Block Models are models of communities. Suppose there are k communities C_1, C_2, \ldots, C_k among a population of n people. Suppose the probability of two people in the same community knowing each other is p and if they are in different communities, the

probability is q (where, q < p).³² We assume the events that person i knows person j are independent across all i and j.

Specifically, we are given an $n \times n$ data matrix A, where, $a_{ij} = 1$ if and only if i and j know each other. We assume the a_{ij} are independent random variables, and use \mathbf{a}_i to denote the i^{th} row of A. It is useful to think of A as the adjacency matrix of a graph, such as the friendship network in Facebook. We will also think of the rows \mathbf{a}_i as data points. The clustering problem is to classify the data points into the communities they belong to. In practice, the graph is fairly sparse, i.e., p and q are small, namely, O(1/n) or $O(\ln n/n)$.

Consider the simple case of two communities with n/2 people in each and with

$$p = \frac{\alpha}{n}$$
; $q = \frac{\beta}{n}$, where, $\alpha, \beta \in O(\ln n)$.

Let **u** and **v** be the centroids of the data points in community one and community two respectively; so, $u_i \approx p$ for $i \in C_1$ and $u_j \approx q$ for $j \in C_2$ and $v_i \approx q$ for $i \in C_1$ and $v_j \approx p$ for $j \in C_2$. We have

$$|\mathbf{u} - \mathbf{v}|^2 \approx \sum_{j=1}^n (u_j - v_j)^2 = \frac{(\alpha - \beta)^2}{n^2} n = \frac{(\alpha - \beta)^2}{n}.$$

Inter-centroid distance $\approx \frac{\alpha - \beta}{\sqrt{n}}.$ (8.1)

On the other hand, the distance between a data point and its cluster centroid is much greater:

For
$$i \in C_1$$
, $E(|\mathbf{a_i} - \mathbf{u}|^2) = \sum_{j=1}^n E((a_{ij} - u_j)^2) = \frac{n}{2}[p(1-p) + q(1-q)] \in \Omega(\alpha + \beta).$

We see that if $\alpha, \beta \in O(\ln n)$, then the ratio

$$\frac{\text{distance between cluster centroids}}{\text{distance of data point to its cluster centroid}} \in O\left(\frac{\sqrt{\ln n}}{\sqrt{n}}\right).$$

So the centroid of a cluster is much farther from data points in its own cluster than it is to the centroid of the other cluster.

Now, consider the k-median objective function. Suppose we wrongly classify a point in C_1 as belonging to C_2 . The extra cost we incur is at most the distance between centroids which is only $O(\sqrt{\ln n}/\sqrt{n})$ times the k-median cost of the data point. So just by examining the cost, we cannot rule out a ε -approximate k-median clustering from misclassifying all points. A similar argument can also be made about k-means objective function.

³²More generally, for each pair of communities a and b, there could be a probability p_{ab} that a person from community a knows a person from community b. But for the discussion here, we take $p_{aa} = p$ for all a and $p_{ab} = q$, for all $a \neq b$.

8.6.2 Gaussian Mixture Model

A second example is the Gaussian Mixture Model with k spherical Gaussians as components, discussed in Section 3.6.2. We saw there that for two Gaussians, each of variance one in each direction, data points are at distance $O(\sqrt{d})$ from their correct centers and if the separation between the centers is O(1), which is much smaller than $O(\sqrt{d})$, an approximate optimal solution could be misclassifying almost all points. We already saw in Section 3.6.2, that SVD of the data matrix A helps in this case.

We will show that a natural and simple SVD-based approach called Spectral Clustering helps solve not only these two examples, but a more general class of problems for which there may or may not be a stochastic model. The result can be qualitatively described by a simple statement when k, the number of clusters, is O(1): If there is some clustering with cluster centroids separated by at least a constant times the "standard deviation", then, we can find a clustering close to this clustering. This should remind the reader of the mnemonic "Means separated by 6 or some constant number of standard deviations".

8.6.3 Standard Deviation without a stochastic model

First, how do we define mean and standard deviation for a clustering problem without assuming a stochastic model of data? In a stochastic model, each cluster consists of independent identically distributed points from a distribution, so the mean of the cluster is just the mean of the distribution. Analogously, we can define the mean as the centroid of the data points in a cluster, whether or not we have a stochastic model. If we had a distribution, it would have a standard deviation in each direction, namely, the square root of the mean squared distance from the mean of the distribution in that direction. But the same definition also applies with no distribution or stochastic model. We give the formal definitions after introducing some notation.

Notation: We will denote by A the data matrix which is $n \times d$, with each row a data point and k will denote the number of clusters. A k-clustering will be represented by a $n \times d$ matrix C; row i of C is the center of the cluster that $\mathbf{a_i}$ belongs to. So, C will have k distinct rows.

The variance of the clustering C along the direction \mathbf{v} , where \mathbf{v} is a vector of length 1, is the mean squared distance of data points from their cluster centers in the direction \mathbf{v} , namely, it is

$$\frac{1}{n}\sum_{i=1}^n \left((\mathbf{a_i} - \mathbf{c_i}) \cdot \mathbf{v} \right)^2.$$

The variance may differ from direction to direction, but we define the variance, denoted σ^2 , of the clustering to be the maximum over all directions, namely,

$$\sigma^{2}(C) = \frac{1}{n} \max_{|\mathbf{v}|=1} \sum_{i=1}^{n} \left((\mathbf{a}_{i} - \mathbf{c}_{i}) \cdot \mathbf{v} \right)^{2} = \frac{1}{n} ||A - C||_{2}^{2}.$$

8.6.4 Spectral Clustering Algorithm

Spectral Clustering - The Algorithm

- 1. Select a value for ϵ .
- Find the top k right singular vectors of data matrix A and let V be the n × k matrix of the top k right singular vectors.
- 3. Select a random row $\mathbf{v_i}$ of V and form a cluster with all rows $\mathbf{v_j}$ such that $|\mathbf{v_i} \mathbf{v_j}| \leq \frac{6k\sigma}{\epsilon}$
- 4. Repeat the above step k times.

Theorem 8.6 If in a k-clustering C, every pair of centers is separated by at least $15k\sigma(C)/\varepsilon$ and every cluster has at least εn points in it, then with probability at least $1-\varepsilon$, **Spectral Clustering** finds a clustering C' which differs from C in at most $\varepsilon^2 n$ points.

Before we prove the theorem, we show that the condition that every pair of cluster centers is separated by $15k\sigma(C)/\varepsilon$ holds for the Stochastic Block Model and Gaussian Mixture models discussed above for appropriate parameter settings. For this, we will need the following theorem from Random Matrix Theory which we do not prove here.

Theorem 8.7 Suppose B is a $n \times d$ matrix with mutually independent, zero mean, random entries with variance ν in $O(\ln n/n)$ that are well-behaved. If $|b_{ij}| \leq 1$ for all i and j or if b_{ij} are Gaussian random variables, they are well-behaved. The theorem works in greater generality. Then, with high probability,

$$||B||_2 \le c\sqrt{n+d}\sqrt{\nu}.$$

Now for the stochastic block model with two communities of n/2 people each and p, q, α , and β as above, we have $E(a_{ij}) = c_{ij}$ with B = A - C:

$$E(b_{ij}) = 0$$
; $\operatorname{var}(b_{ij}) = p(1-p)$ or $q(1-q) \le p$.

Setting $\nu = p$, the theorem gives

$$||A - C||_2 \le c\sqrt{n}\sqrt{p} = c\sqrt{\alpha} \implies \sigma(C) \le \frac{\sqrt{\alpha}}{\sqrt{n}}.$$

From (8.1) inter-center separation is $\frac{\alpha-\beta}{\sqrt{n}}$. Thus, the condition of the theorem is satisfied as long as $\sqrt{\alpha} \in \Omega(\alpha - \beta)$, which is a reasonable assumption in the regime when α is at least a large constant.

The proof for the Gaussian mixture model is similar. Suppose we have a mixture of k Gaussians and A is a data matrix with n independent, identically distributed samples from the mixture as its rows. The Gaussians need not be spherical. Let σ_{max} be the maximum standard deviation of any of the k Gaussians in any direction. We again consider C to

be made up of the means of the Gaussians. Now the theorem is satisfied by A - C with $\nu = \sigma_{\max}^2$. For $k \in O(1)$, it is easy to see that the hypothesis of Theorem 8.6 is satisfied provided the means of the component Gaussians are separated by $\Omega(\sigma_{\max})$.

The proof of the Theorem 8.6 relies on a crucial lemma, which is simple to prove.

Lemma 8.8 Suppose A is an $n \times d$ matrix and suppose V is obtained by projecting the rows of A to the subspace of the first k right singular vectors of A. For any matrix C of rank less than or equal to k

$$||V - C||_F^2 \le 8k||A - C||_2^2.$$

Note: V is just one matrix. But it is close to every C, in the sense $||V - C||_F^2 \leq 8k||A - C||_2^2$. While this seems contradictory, the point of the lemma is that for C far away from V, $||A - C||_2$ will be high.

Proof: Since the rank of (V - C) is less than or equal to 2k,

$$||V - C||_F^2 \le 2k||V - C||_2^2 \text{ and}$$
$$||V - C||_2 \le ||V - A||_2 + ||A - C||_2 \le 2||A - C||_2.$$

The last inequality follows since V is the best rank k approximation in spectral norm and C has rank at most k. The lemma follows.

Proof of Theorem 8.6: We use the lemma to argue that barring a few exceptions, most $\bar{\mathbf{a}}_{\mathbf{i}}$ are at distance at most $3k\sigma(C)/\varepsilon$ to the corresponding $\mathbf{c}_{\mathbf{i}}$. This will imply that for most i, the point $\bar{\mathbf{a}}_{\mathbf{i}}$ will be close distance at most to most $6k\sigma(C)/\varepsilon$ to most other $\bar{\mathbf{a}}_{\mathbf{j}}$ in its own cluster. Since we assumed the cluster centers of C are well separated this will imply that for most i and j in different clusters, $|\mathbf{v}_{\mathbf{i}} - \mathbf{v}_{\mathbf{j}}| \geq 9k\sigma(C)/\varepsilon$. This will enable us to prove that the distance based clustering step, Step 3, of the algorithm works.

Define M to be the set of exceptions:

$$M = \{ i : |\mathbf{ab}_{\mathbf{i}} - \mathbf{c}_{\mathbf{i}}| \ge 3k\sigma(C)/\varepsilon \}.$$

Since $||V - C||_F^2 = \sum_i |\mathbf{v_i} - \mathbf{c_i}|^2 \ge \sum_{i \in M} |\mathbf{v_i} - \mathbf{c_i}|^2 \ge |M| \frac{9k^2 \sigma^2(C)}{\varepsilon^2}$, using the lemma we get:

$$|M|\frac{9k^2\sigma^2(C)}{\varepsilon^2} \le ||V - C||_F^2 \le 8kn\sigma^2(C) \implies |M| \le \frac{8\varepsilon^2 n}{9k}.$$
(8.2)

For
$$i, j \notin M$$
, i, j in the same cluster in C ,
 $|\mathbf{v_i} - \mathbf{v_j}| \le |\mathbf{v_i} - \mathbf{c_i}| + |\mathbf{c_i} - \mathbf{v_j}| \le \frac{6k\sigma(C)}{\varepsilon}$. (8.3)

But for $l \neq l'$, by hypothesis the cluster centers are at least $15k\sigma(C)/\varepsilon$ apart. This implies that

For
$$i, k \notin M$$
, i, k in different clusters in C

$$|\mathbf{v}_{i} - \mathbf{v}_{k}| \ge |\mathbf{c}_{i} - \mathbf{c}_{k}| - |\mathbf{c}_{i} - \mathbf{v}_{i}| - |\mathbf{v}_{k} - \mathbf{c}_{k}| \ge \frac{9k\sigma(C)}{2\varepsilon}.$$
(8.4)

We will show by induction on the number of iterations of Step 3 the invariant that at the end of t iterations of Step 3, S consists of the union of k - t of the $k \ C_{\ell} \setminus M$ plus a subset of M. In other words, but for elements of M, S is precisely the union of k - t clusters of C. Clearly, this holds for t = 0. Suppose it holds for a t. Suppose in iteration t + 1of Step 3 of the algorithm, we choose an $i_0 \notin M$ and say i_0 is in cluster ℓ in C. Then by (8.3) and (8.4), T will contain all points of $C_{\ell} \setminus M$ and will contain no points of $C_{\ell'} \setminus M$ for any $\ell' \neq \ell$. This proves the invariant still holds after the iteration. Also, the cluster returned by the algorithm will agree with C_{ℓ} except possibly on M provided $i_0 \notin M$.

Now by (8.2), $|M| \leq \varepsilon^2 n$ and since each $|C_\ell| \geq \varepsilon n$, we have that $|C_\ell \setminus M| \geq (\varepsilon - \varepsilon^2)n$. If we have done less than k iterations of Step 3 and not yet peeled off C_ℓ , then there are still $(\varepsilon - \varepsilon^2)n$ points of $C_\ell \setminus M$ left. So the probability that the next pick i_0 will be in M is at most $|M|/(\varepsilon - \varepsilon^2)n \leq \varepsilon/k$ by (8.2). So with probability at least $1 - \varepsilon$ all the k i_0 's we pick are out of M and the theorem follows.

8.7 High-Density Clusters

We now turn from the assumption that clusters are center-based to the assumption that clusters consist of high-density regions, separated by low-density moats such as in Figure 8.1.

8.7.1 Single-linkage

One natural algorithm for clustering under the high-density assumption is called *single* linkage. This algorithm begins with each point in its own cluster and then repeatedly merges the two "closest" clusters into one, where the distance between two clusters is defined as the minimum distance between points in each cluster. That is, $d_{min}(C, C') = \min_{\mathbf{x} \in C, \mathbf{y} \in C'} d(\mathbf{x}, \mathbf{y})$, and the algorithm merges the two clusters C and C' whose d_{min} value is smallest over all pairs of clusters breaking ties arbitrarily. It then continues until there are only k clusters. This is called an agglomerative clustering algorithm because it begins with many clusters and then starts merging, or agglomerating them together.³³ Singlelinkage is equivalent to running Kruskal's minimum-spanning-tree algorithm, but halting when there are k trees remaining. The following theorem is fairly immediate.

³³Other agglomerative algorithms include *complete linkage* which merges the two clusters whose *maximum* distance between points is smallest, and Ward's algorithm described earlier that merges the two clusters that cause the *k*-means cost to increase by the least.

Theorem 8.9 Suppose the desired clustering C_1^*, \ldots, C_k^* satisfies the property that there exists some distance σ such that

- 1. any two data points in different clusters have distance at least σ , and
- 2. for any cluster C_i^* and any partition of C_i^* into two non-empty sets A and $C_i^* \setminus A$, there exist points on each side of the partition of distance less than σ .

Then, single-linkage will correctly recover the clustering C_1^*, \ldots, C_k^* .

Proof: Consider running the algorithm until all pairs of clusters C and C' have $d_{min}(C, C') \ge \sigma$. At that point, by (b), each target cluster C_i^* will be fully contained within some cluster of the single-linkage algorithm. On the other hand, by (a) and by induction, each cluster C of the single-linkage algorithm will be fully contained within some C_i^* of the target clustering, since any merger of subsets of distinct target clusters would require $d_{min} \ge \sigma$. Therefore, the single-linkage clusters are indeed the target clusters.

8.7.2 Robust linkage

The single-linkage algorithm is fairly brittle. A few points bridging the gap between two different clusters can cause it to do the wrong thing. As a result, there has been significant work developing more robust versions of the algorithm.

One commonly used robust version of single linkage is Wishart's algorithm. We can view single-linkage as growing balls of radius r around each datapoint, starting with r = 0and then gradually increasing r, connecting two points when the balls around them touch. The clusters are the connected components of this graph. To address the issue of a few points causing an incorrect merger, Wishart's algorithm has a parameter t, and only considers a point to be live if its ball of radius r contains at least t points. It then only makes a connection between live points. The idea is that if t is modestly large, then a thin string of points between two dense clusters will not cause a spurious merger.

In fact, if one slightly modifies the algorithm to define a point to be live if its ball of radius r/2 contains at least t points, then it is known [?] that a value of $t = O(d \log n)$ is sufficient to recover a nearly correct solution under a natural distributional formulation of the clustering problem. Specifically, suppose data points are drawn from some probability distribution D over \mathbb{R}^d , and that the clusters correspond to high-density regions surrounded by lower-density moats. More specifically, the assumption is that

- 1. for some distance $\sigma > 0$, the σ -interior of each target cluster C_i^* has density at least some quantity λ (the σ -interior is the set of all points at distance at least σ from the boundary of the cluster),
- 2. the region between target clusters has density less than $\lambda(1-\epsilon)$ for some $\epsilon > 0$,
- 3. the clusters should be separated by distance greater than 2σ , and

4. the σ -interior of the clusters contains most of their probability mass.

Then, for sufficiently large n, the algorithm will with high probability find nearly correct clusters. In this formulation, we allow points in low-density regions that are not in any target clusters at all. For details, see [?].

Robust Median Neighborhood Linkage robustifies single linkage in a different way. This algorithm guarantees that if it is possible to delete a small fraction of the data such that for all remaining points x, most of their $|C^*(x)|$ nearest neighbors indeed belong to their own cluster $C^*(x)$, then the hierarchy on clusters produced by the algorithm will include a close approximation to the true clustering. We refer the reader to [?] for the algorithm and proof.

8.8 Kernel Methods

Kernel methods combine aspects of both center-based and density-based clustering. In center-based approaches like k-means or k-center, once the cluster centers are fixed, the Voronoi diagram of the cluster centers determines which cluster each data point belongs to. This implies that clusters are pairwise linearly separable.

If we believe that the true desired clusters may not be linearly separable, and yet we wish to use a center-based method, then one approach, as in the chapter on learning, is to use a kernel. Recall that a kernel function $K(\mathbf{x}, \mathbf{y})$ can be viewed as performing an implicit mapping ϕ of the data into a possibly much higher dimensional space, and then taking a dot-product in that space. That is, $K(\mathbf{x}, \mathbf{y}) = \phi(\mathbf{x}) \cdot \phi(\mathbf{y})$. This is then viewed as the affinity between points \mathbf{x} and \mathbf{y} . We can extract distances in this new space using the equation $|\mathbf{z_1} - \mathbf{z_2}|^2 = \mathbf{z_1} \cdot \mathbf{z_1} + \mathbf{z_2} \cdot \mathbf{z_2} - 2\mathbf{z_1} \cdot \mathbf{z_2}$, so in particular we have $|\phi(\mathbf{x}) - \phi(\mathbf{y})|^2 = K(\mathbf{x}, \mathbf{x}) + K(\mathbf{y}, \mathbf{y}) - 2K(\mathbf{x}, \mathbf{y})$. We can then run a center-based clustering algorithm on these new distances.

One popular kernel function to use is the Gaussian kernel. The Gaussian kernel uses an affinity measure that emphasizes closeness of points and drops off exponentially as the points get farther apart. Specifically, we define the affinity between points \mathbf{x} and \mathbf{y} by

$$K(\mathbf{x}, \mathbf{y}) = e^{-\frac{1}{2\sigma^2} \|\mathbf{x} - \mathbf{y}\|^2}.$$

Another way to use affinities is to put them in an affinity matrix, or weighted graph. This graph can then be separated into clusters using a graph partitioning procedure such as in the following section.

8.9 Recursive Clustering based on Sparse cuts

We now consider the case that data are nodes in an undirected connected graph G(V, E) where an edge indicates that the end point vertices are similar. Recursive clustering starts with all vertices in one cluster and recursively splits a cluster into two parts

whenever there is a small number of edges from one part to the other part of the cluster. Formally, for two disjoint sets S and T of vertices, define

$$\Phi(S,T) = \frac{\text{Number of edges from } S \text{ to } T}{\text{Total number of edges incident to } S \text{ in } G}.$$

 $\Phi(S,T)$ measures the relative strength of similarities between S and T. Let d(i) be the degree of vertex i and for a subset S of vertices, let $d(S) = \sum_{i \in S} d(i)$. Let m be the total number of edges. The following algorithm aims to cut only a small fraction of the edges and to produce clusters that are internally consistent in that no subset of the cluster has low similarity to the rest of the cluster.

Recursive Clustering: Select an appropriate value for ϵ . If a current cluster W has a subset S with $d(S) \leq \frac{1}{2}d(W)$ and $\Phi(S,T) \leq \epsilon$, then split W into two clusters S and $W \setminus S$. Repeat until no such split is possible.

Theorem 8.10 At termination of **Recursive Clustering**, the total number of edges between vertices in different clusters is at most $O(\varepsilon m \ln n)$.

Proof: Each edge between two different clusters at the end was "cut up" at some stage by the algorithm. We will "charge" edge cuts to vertices and bound the total charge. When the algorithm partitions a cluster W into S and $W \setminus S$ with $d(S) \leq (1/2)d(W)$, each $k \in S$ is charged $\frac{d(k)}{d(W)}$ times the number of edges being cut. Since $\Phi(S, W \setminus S) \leq \varepsilon$, the charge added to each $k \in W$ is a most $\varepsilon d(k)$. A vertex is charged only when it is in the smaller part $(d(S) \leq d(W)/2)$ of the cut. So between any two times it is charged, d(W) is reduced by a factor of at least two and so a vertex can be charged at most $\log_2 m \leq O(\ln n)$ times, proving the Theorem.

Implementing the algorithm requires computing $\operatorname{Min}_{S \subseteq W} \Phi(S, W \setminus S)$ which is an NPhard problem. So the theorem cannot be implemented right away. Luckily, eigenvalues and eigenvectors, which can be computed fast, give an approximate answer. The connection between eigenvalues and sparsity, known as Cheeger's inequality, is deep with applications to Markov chains among others. We do not discuss this here.

8.10 Dense Submatrices and Communities

Represent n data points in d-space by the rows of an $n \times d$ matrix A. Assume that A has all nonnegative entries. Examples to keep in mind for this section are the document-term matrix and the customer-product matrix. We address the question of how to define and find efficiently a coherent large subset of rows. To this end, the matrix A can be represented by a bipartite graph. One side has a vertex for each row and the other side a vertex for each column. Between the vertex for row i and the vertex for column j, there is an edge with weight a_{ij} .

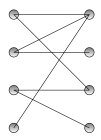


Figure 8.4: Example of a bipartite graph.

We want a subset S of row vertices and a subset T of column vertices so that

$$A(S,T) = \sum_{i \in S, j \in T} a_{ij}$$

is high. This simple definition is not good since A(S,T) will be maximized by taking all rows and columns. We need a balancing criterion that ensures that A(S,T) is high relative to the sizes of S and T. One possibility is to maximize $\frac{A(S,T)}{|S||T|}$. This is not a good measure either, since it is maximized by the single edge of highest weight. The definition we use is the following. Let A be a matrix with nonnegative entries. For a subset S of rows and a subset T of columns, the *density* d(S,T) of S and T is $d(S,T) = \frac{A(S,T)}{\sqrt{|S||T|}}$. The *density* d(A) of A is defined as the maximum value of d(S,T) over all subsets of rows and columns. This definition applies to bipartite as well as non bipartite graphs.

One important case is when A's rows and columns both represent the same set and a_{ij} is the similarity between object i and object j. Here $d(S, S) = \frac{A(S,S)}{|S|}$. If A is an $n \times n$ 0-1 matrix, it can be thought of as the adjacency matrix of an undirected graph, and d(S, S) is the average degree of a vertex in S. The subgraph of maximum average degree in a graph can be found exactly by network flow techniques, as we will show in the next section. We do not know an efficient (polynomial-time) algorithm for finding d(A) exactly in general. However, we show that d(A) is within a $O(\log^2 n)$ factor of the top singular value of A assuming $|a_{ij}| \leq 1$ for all i and j. This is a theoretical result. The gap may be much less than $O(\log^2 n)$ for many problems, making singular values and singular vectors quite useful. Also, S and T with $d(S, T) \geq \Omega(d(A)/\log^2 n)$ can be found algorithmically.

Theorem 8.11 Let A be an $n \times d$ matrix with entries between 0 and 1. Then

$$\sigma_1(A) \ge d(A) \ge \frac{\sigma_1(A)}{4\log n \log d}.$$

Furthermore, subsets S and T satisfying $d(S,T) \geq \frac{\sigma_1(A)}{4 \log n \log d}$ may be found from the top singular vector of A.

Proof: Let S and T be the subsets of rows and columns that achieve d(A) = d(S, T). Consider an *n*-vector **u** which is $\frac{1}{\sqrt{|S|}}$ on S and 0 elsewhere and a *d*-vector **v** which is $\frac{1}{\sqrt{|T|}}$ on T and 0 elsewhere. Then,

$$\sigma_1(A) \ge \mathbf{u}^T A \mathbf{v} = \sum_{ij} u_i v_j a_{ij} = d(S, T) = d(A)$$

establishing the first inequality.

To prove the second inequality, express $\sigma_1(A)$ in terms of the first left and right singular vectors \mathbf{x} and \mathbf{y} .

$$\sigma_1(A) = \mathbf{x}^T A \mathbf{y} = \sum_{i,j} x_i a_{ij} y_j, \qquad |\mathbf{x}| = |\mathbf{y}| = 1.$$

Since the entries of A are nonnegative, the components of the first left and right singular vectors must all be nonnegative, that is, $x_i \ge 0$ and $y_j \ge 0$ for all i and j. To bound $\sum_{i,j} x_i a_{ij} y_j$, break the summation into $O(\log n \log d)$ parts. Each part corresponds to a given α and β and consists of all i such that $\alpha \le x_i < 2\alpha$ and all j such that $\beta \le y_i < 2\beta$. The $\log n \log d$ parts are defined by breaking the rows into $\log n$ blocks with α equal to $\frac{1}{2} \frac{1}{\sqrt{n}}, \frac{1}{\sqrt{n}}, \frac{2}{\sqrt{d}}, \frac{4}{\sqrt{n}}, \ldots, 1$ and by breaking the columns into $\log d$ blocks with β equal to $\frac{1}{2} \frac{1}{\sqrt{d}}, \frac{1}{\sqrt{d}}, \frac{2}{\sqrt{d}}, \frac{4}{\sqrt{d}}, \ldots, 1$. The i such that $x_i < \frac{1}{2\sqrt{n}}$ and the j such that $y_j < \frac{1}{2\sqrt{d}}$ will be ignored at a loss of at most $\frac{1}{4}\sigma_1(A)$. Exercise (8.28) proves the loss is at most this amount.

Since $\sum_{i} x_i^2 = 1$, the set $S = \{i | \alpha \leq x_i < 2\alpha\}$ has $|S| \leq \frac{1}{\alpha^2}$ and similarly, $T = \{j | \beta \leq y_j \leq 2\beta\}$ has $|T| \leq \frac{1}{\beta^2}$. Thus

$$\sum_{\substack{i \\ \alpha \leq x_i \leq 2\alpha}} \sum_{\substack{j \\ \beta \leq y_j \leq 2\beta}} x_i y_j a_{ij} \leq 4\alpha \beta A(S,T)$$
$$\leq 4\alpha \beta d(S,T) \sqrt{|S||T|}$$
$$\leq 4d(S,T)$$
$$< 4d(A).$$

From this it follows that

$$\sigma_1(A) \le 4d(A)\log n\log d$$

or

$$d(A) \ge \frac{\sigma_1(A)}{4\log n \log d}$$

proving the second inequality.

It is clear that for each of the values of (α, β) , we can compute A(S, T) and d(S, T) as above and taking the best of these d(S, T) 's gives us an algorithm as claimed in the theorem.

Note that in many cases, the nonzero values of x_i and y_j after zeroing out the low entries will only go from $\frac{1}{2}\frac{1}{\sqrt{n}}$ to $\frac{c}{\sqrt{n}}$ for x_i and $\frac{1}{2}\frac{1}{\sqrt{d}}$ to $\frac{c}{\sqrt{d}}$ for y_j , since the singular vectors

are likely to be balanced given that a_{ij} are all between 0 and 1. In this case, there will be O(1) groups only and the log factors disappear.

Another measure of density is based on similarities. Recall that the similarity between objects represented by vectors (rows of A) is defined by their dot products. Thus, similarities are entries of the matrix AA^T . Define the average cohesion f(S) of a set S of rows of A to be the sum of all pairwise dot products of rows in S divided by |S|. The average cohesion of A is the maximum over all subsets of rows of the average cohesion of the subset.

Since the singular values of AA^T are squares of singular values of A, we expect f(A) to be related to $\sigma_1(A)^2$ and $d(A)^2$. Indeed it is. We state the following without proof.

Lemma 8.12 $d(A)^2 \le f(A) \le d(A) \log n$. Also, $\sigma_1(A)^2 \ge f(A) \ge \frac{c\sigma_1(A)^2}{\log n}$.

f(A) can be found exactly using flow techniques as we will see later.

8.11 Community Finding and Graph Partitioning

Assume that data are nodes in a possibly weighted graph where edges represent some notion of affinity between their endpoints. In particular, let G = (V, E) be a weighted graph. Given two sets of nodes S and T, define

$$E(S,T) = \sum_{\substack{i \in S \\ j \in T}} e_{ij}.$$

We then define the *density* of a set S to be

$$d(S,S) = \frac{E(S,S)}{|S|}.$$

If G is an undirected graph, then d(S, S) can be viewed as the average degree in the vertex-induced subgraph over S. The set S of maximum density is therefore the subgraph of maximum average degree. Finding such a set can be viewed as finding a tight-knit community inside some network. In the next section, we describe an algorithm for finding such a set using network flow techniques.

8.11.1 Flow Methods

Here we consider dense induced subgraphs of a graph. An induced subgraph of a graph consisting of a subset of the vertices of the graph along with all edges of the graph that connect pairs of vertices in the subset of vertices. We show that finding an induced subgraph with maximum average degree can be done by network flow techniques. This is simply maximizing the density d(S, S) over all subsets S of the graph. First consider the problem of finding a subset of vertices such that the induced subgraph has average

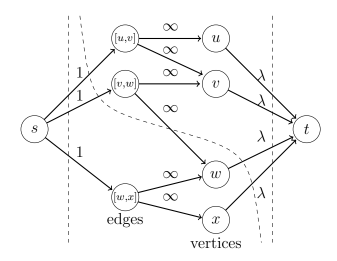


Figure 8.5: The directed graph H used by the flow technique to find a dense subgraph

degree at least λ for some parameter λ . Then do a binary search on the value of λ until the maximum λ for which there exists a subgraph with average degree at least λ is found.

Given a graph G in which one wants to find a dense subgraph, construct a directed graph H from the given graph and then carry out a flow computation on H. H has a node for each edge of the original graph, a node for each vertex of the original graph, plus two additional nodes s and t. There is a directed edge with capacity one from s to each node corresponding to an edge of the original graph and a directed edge with infinite capacity from each node corresponding to an edge of the original graph to the two nodes corresponding to the vertices the edge connects. Finally, there is a directed edge with capacity λ from each node corresponding to a vertex of the original graph to t.

Notice there are three types of cut sets of the directed graph that have finite capacity. The first cuts all arcs from the source. It has capacity e, the number of edges of the original graph. The second cuts all edges into the sink. It has capacity λv , where v is the number of vertices of the original graph. The third cuts some arcs from s and some arcs into t. It partitions the set of vertices and the set of edges of the original graph into two blocks. The first block contains the source node s, a subset of the edges e_s , and a subset of the vertices v_s defined by the subset of edges. The first block must contain both end points of each edge in e_s ; otherwise an infinite arc will be in the cut. The second block contains t and the remaining edges and vertices. The edges in this second block either connect vertices in the second block or have one endpoint in each block. The cut set will cut some infinite arcs from edges not in e_s coming into vertices in v_s . However, these arcs are directed from nodes in the block containing t to nodes in the block containing s. Note that any finite capacity cut that leaves an edge node connected to s must cut the two related vertex nodes from t. Thus, there is a cut of capacity $e - e_s + \lambda v_s$ where v_s and e_s are the vertices and edges of a subgraph. For this cut to be the minimal cut, the

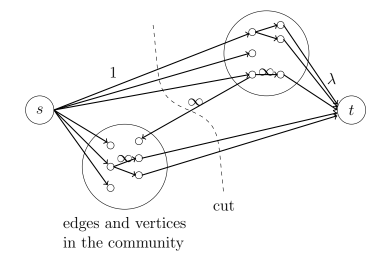


Figure 8.6: Cut in flow graph

quantity $e - e_s + \lambda v_s$ must be minimal over all subsets of vertices of the original graph and the capcity must be less than e and also less than λv .

If there is a subgraph with v_s vertices and e_s edges where the ratio $\frac{e_s}{v_s}$ is sufficiently large so that $\frac{e_s}{v_s} > \frac{e}{v}$, then for λ such that $\frac{e_s}{v_s} > \lambda > \frac{e}{v}$, $e_s - \lambda v_s > 0$ and $e - e_s + \lambda v_s < e$. Similarly $e < \lambda v$ and thus $e - e_s + \lambda v_s < \lambda v$. This implies that the cut $e - e_s + \lambda v_s$ is less than either e or λv and the flow algorithm will find a nontrivial cut and hence a proper subset. For different values of λ in the above range there maybe different nontrivial cuts.

Note that for a given density of edges, the number of edges grows as the square of the number of vertices and $\frac{e_s}{v_s}$ is less likely to exceed $\frac{e}{v}$ if v_S is small. Thus, the flow method works well in finding large subsets since it works with $\frac{e_S}{v_S}$. To find small communities one would need to use a method that worked with $\frac{e_S}{v_S^2}$ as the following example illustrates.

Example: Consider finding a dense subgraph of 1,000 vertices and 2,000 internal edges in a graph of 10^6 vertices and 6×10^6 edges. For concreteness, assume the graph was generated by the following process. First, a 1,000-vertex graph with 2,000 edges was generated as a random regular degree four graph. The 1,000-vertex graph was then augmented to have 10^6 vertices and edges were added at random until all vertices were of degree 12. Note that each vertex among the first 1,000 has four edges to other vertices among the first 1,000 has four edges to other vertices is much denser than the whole graph in some sense. Although the subgraph induced by the 1,000 vertices has four edges per vertex and the full graph has twelve edges per vertex, the probability of two vertices of the 1,000 being connected by an edge is much higher than for the graph as a whole. The probability is given by the ratio of the actual number of edges connecting vertices among the 1,000 to the number of possible edges if the vertices formed a complete

graph.

$$p = \frac{e}{\left(\binom{v}{2}\right)} = \frac{2e}{v(v-1)}$$

For the 1,000 vertices, this number is $p = \frac{2 \times 2,000}{1,000 \times 999} \cong 4 \times 10^{-3}$. For the entire graph this number is $p = \frac{2 \times 6 \times 10^6}{10^6 \times 10^6} = 12 \times 10^{-6}$. This difference in probability of two vertices being connected should allow us to find the dense subgraph.

In our example, the cut of all arcs out of s is of capacity 6×10^6 , the total number of edges in the graph, and the cut of all arcs into t is of capacity λ times the number of vertices or $\lambda \times 10^6$. A cut separating the 1,000 vertices and 2,000 edges would have capacity $6 \times 10^6 - 2,000 + \lambda \times 1,000$. This cut cannot be the minimum cut for any value of λ since $\frac{e_s}{v_s} = 2$ and $\frac{e}{v} = 6$, hence $\frac{e_s}{v_s} < \frac{e}{v}$. The point is that to find the 1,000 vertices, we have to maximize $A(S,S)/|S|^2$ rather than A(S,S)/|S|. Note that $A(S,S)/|S|^2$ penalizes large |S| much more and therefore can find the 1,000 node "dense" subgraph.

8.12 Axioms for Clustering

Each clustering algorithm tries to optimize some criterion, like the sum of squared distances to the nearest cluster center, over all possible clusterings. We have seen many different optimization criteria in this chapter and many more are used. Now, we take a step back and look at properties that one might want a clustering criterion or algorithm to have, and ask which criteria have them and which sets of properties are simultaneously achievable by some algorithm? We begin with a negative statement. We present three seemingly desirable properties of a clustering algorithm and then show that no algorithm can satisfy them simultaneously. Next we argue that these requirements are too stringent and under more reasonable requirements, a slightly modified form of the sum of Euclidean distance squared between all pairs of points inside the same cluster is indeed a measure satisfying the desired properties.

8.12.1 An Impossibility Result

Let $\mathcal{A}(d)$ denote the clustering found by clustering algorithm \mathcal{A} using the distance function d on a set S. The clusters of the clustering $\mathcal{A}(d)$ form a partition Γ of S.

The first property we consider of a clustering algorithm is scale invariance. A clustering algorithm \mathcal{A} is scale invariant if for any $\alpha > 0$, $\mathcal{A}(d) = \mathcal{A}(\alpha d)$. That is, multiplying all distances by some scale factor does not change the algorithm's clustering.

A clustering algorithm \mathcal{A} is *rich* (full/complete) if for every partitioning Γ of S there exists a distance function d over S such that $\mathcal{A}(d) = \Gamma$. That is, for any desired partitioning, we can find a set of distances so that the clustering algorithm returns the desired partitioning.

A clustering algorithm is *consistent* if increasing the distance between points in different clusters and reducing the distance between points in the same cluster does not change the clusters produced by the clustering algorithm.

We now show that no clustering algorithm \mathcal{A} can satisfy all three of scale invariance, richness, and consistency.³⁴

Theorem 8.13 No algorithm \mathcal{A} can satisfy all three of scale invariance, richness, and consistency.

Proof: Let's begin with the simple case of just two points, $S = \{a, b\}$. By richness there must be some distance d(a, b) such that \mathcal{A} produces two clusters and some other distance d'(a, b) such that \mathcal{A} produces one cluster. But this then violates scale-invariance.

Let us now turn to the general case of n points, $S = \{a_1, \ldots, a_n\}$. By richness, there must exist some distance function d such that \mathcal{A} puts all of S into a single cluster, and some other distance function d' such that \mathcal{A} puts each point of S into its own cluster. Let ϵ be the minimum distance between points in d, and let Δ be the maximum distance between points in d'. Define $d'' = \alpha d'$ for $\alpha = \epsilon/\Delta$; i.e., uniformly shrink distances in d'until they are all less than or equal to the minimum distance in d. By scale invariance, $\mathcal{A}(d'') = \mathcal{A}(d')$, so under d'', \mathcal{A} puts each point of S into its own cluster. However, note that for each pair of points a_i, a_j we have $d''(a_i, a_j) \leq d(a_i, a_j)$. This means we can reach d'' from d by just reducing distances between points in the same cluster (since all points are in the same cluster under d). So, the fact that \mathcal{A} behaves differently on d'' and dviolates consistency.

8.12.2 Satisfying two of three

There exist natural clustering algorithms satisfying any two of the three axioms. For example, different versions of the *single linkage* algorithm described in Section 8.7 satisfy different two of the three conditions.

Theorem 8.14

- 1. The single linkage clustering algorithm with the k-cluster stopping condition (stop when there are k clusters), satisfies scale-invariance and consistency. We do not get richness since we only get clusterings with k clusters.
- 2. The single linkage clustering algorithm with scale α stopping condition satisfies scale invariance and richness. The scale α stopping condition is to stop when the closest pair of clusters is of distance greater than or equal to αd_{\max} where d_{\max} is the maximum pair wise distance. Here we do not get consistency. If we select one distance

 $^{^{34}}$ A technical point here: we do not allow d to have distance 0 between two distinct points of S. Else, a simple algorithm that satisfies all three properties is simply "place two points into the same cluster if they have distance 0, else place them into different clusters".



Figure 8.7: Illustration of the objection to the consistency axiom. Reducing distances between points in a cluster may suggest that the cluster be split into two.

between clusters and increase it significantly until it becomes d_{\max} and in addition αd_{\max} exceeds all other distances, the resulting clustering has just one cluster containing all of the points.

3. The single linkage clustering algorithm with the distance r stopping condition, stop when the inter-cluster distances are all at least r, satisfies richness and consistency; but not scale invariance.

Proof: (1) Scale-invariance is easy to see. If one scales up all distances by a factor, then at each point in the algorithm, the same pair of clusters will be closest. The argument for consistency is more subtle. Since edges inside clusters of the final clustering can only be decreased and since edges between clusters can only be increased, the edges that led to merges between any two clusters are less than any edge between the final clusters. Since the final number of clusters is fixed, these same edges will cause the same merges unless the merge has already occurred due to some other edge that was inside a final cluster having been shortened even more. No edge between two final clusters can cause a merge before all the above edges have been considered. At this time the final number of clusters has been reached and the process of merging has stopped. Parts (2) and (3) are straightforward.

Note that one may question both the consistency axiom and the richness axiom. The following are two possible objections to the consistency axiom. Consider the two clusters in Figure 8.7. If one reduces the distance between points in cluster B, they might get an arrangement that should be three clusters instead of two.

The other objection, which applies to both the consistency and the richness axioms, is that they force many unrealizable distances to exist. For example, suppose the points were in Euclidean d space and distances were Euclidean. Then, there are only nd degrees of freedom. But the abstract distances used here have $O(n^2)$ degrees of freedom since the distances between the $O(n^2)$ pairs of points can be specified arbitrarily. Unless d is about n, the abstract distances are too general. The objection to richness is similar. If for npoints in Euclidean d space, the clusters are formed by hyper planes each cluster may be a Voronoi cell or some other polytope, then as we saw in the theory of VC dimensions Section ?? there are only $\binom{n}{d}$ interesting hyper planes each defined by d of the n points. If k clusters are defined by bisecting hyper planes of pairs of points, there are only n^{dk^2} possible clustering's rather than the 2^n demanded by richness. If d and k are significantly less than n, then richness is not reasonable to demand. In the next section, we will see a possibility result to contrast with this impossibility theorem.

8.12.3 Relaxing the axioms

Given that no clustering algorithm can satisfy scale invariance, richness, and consistency, one might want to relax the axioms in some way. Then one gets the following results.

1. Single linkage with a distance stopping condition satisfies a relaxed scale-invariance property that states that for $\alpha > 1$, then $f(\alpha d)$ is a refinement of f(d).

2. Define refinement consistency to be that shrinking distances within a cluster or expanding distances between clusters gives a refinement of the clustering. Single linkage with α stopping condition satisfies scale invariance, refinement consistency and richness except for the trivial clustering of all singletons.

8.12.4 A Satisfiable Set of Axioms

In this section, we propose a different set of axioms that are reasonable for distances between points in Euclidean space and show that the clustering measure, the sum of squared distances between all pairs of points in the same cluster, slightly modified, is consistent with the new axioms. We assume through the section that points are in Euclidean *d*-space. Our three new axioms follow.

We say that a clustering algorithm satisfies the *consistency condition* if, for the clustering produced by the algorithm on a set of points, moving a point so that its distance to any point in its own cluster is not increased and its distance to any point in a different cluster is not decreased, then the algorithm returns the same clustering after the move.

Remark: Although it is not needed in the sequel, it is easy to see that for an infinitesimal perturbation dx of x, the perturbation is consistent if and only if each point in the cluster containing x lies in the half space through x with dx as the normal and each point in a different cluster lies in the other half space.

An algorithm is *scale-invariant* if multiplying all distances by a positive constant does not change the clustering returned.

An algorithm has the *richness* property if for any set K of k distinct points in the ambient space, there is some placement of a set S of n points to be clustered so that the algorithm returns a clustering with the points in K as centers. So there are k clusters, each cluster consisting of all points of S closest to one particular point of K.

We will show that the following algorithm satisfies these three axioms.

Balanced *k*-means algorithm

Among all partitions of the input set of n points into k sets, each of size n/k, return the one that minimizes the sum of squared distances between all pairs of points in the same cluster.

Theorem 8.15 The balanced k-means algorithm satisfies the consistency condition, scale invariance, and the richness property.

Proof: Scale invariance is obvious. Richness is also easy to see. Just place n/k points of S to coincide with each point of K. To prove consistency, define the *cost* of a cluster T to be the sum of squared distances of all pairs of points in T.

Suppose S_1, S_2, \ldots, S_k is an optimal clustering of S according to the balanced kmeans algorithm. Move a point $x \in S_1$ to z so that its distance to each point in S_1 is non increasing and its distance to each point in S_2, S_3, \ldots, S_k is non decreasing. Suppose T_1, T_2, \ldots, T_k is an optimal clustering after the move. Without loss of generality assume $z \in T_1$. Define $\tilde{T}_1 = (T_1 \setminus \{z\}) \cup \{x\}$ and $\tilde{S}_1 = (S_1 \setminus \{x\}) \cup \{z\}$. Note that $\tilde{T}_1, T_2, \ldots, T_k$ is a clustering before the move, although not necessarily an optimal clustering. Thus

$$\cot\left(\tilde{T}_{1}\right) + \cot\left(T_{2}\right) + \dots + \cot\left(T_{k}\right) \ge \cot\left(S_{1}\right) + \cot\left(S_{2}\right) + \dots + \cot\left(S_{k}\right).$$

If $\operatorname{cost}(T_1) - \operatorname{cost}(\tilde{T}_1) \ge \operatorname{cost}(\tilde{S}_1) - \operatorname{cost}(S_1)$ then

$$\operatorname{cost}(T_1) + \operatorname{cost}(T_2) + \dots + \operatorname{cost}(T_k) \ge \operatorname{cost}\left(\tilde{S}_1\right) + \operatorname{cost}\left(S_2\right) + \dots + \operatorname{cost}\left(S_k\right).$$

Since T_1, T_2, \ldots, T_k is an optimal clustering after the move, so also must be $\tilde{S}_1, S_2, \ldots, S_k$ proving the theorem.

It remains to show that $\cot(T_1) - \cot(\tilde{T}_1) \ge \cot(\tilde{S}_1) - \cot(S_1)$. Let u and v stand for elements other than x and z in S_1 and T_1 . The terms $|u-v|^2$ are common to T_1 and \tilde{T}_1 on the left hand side and cancel out. So too on the right hand side. So we need only prove

$$\sum_{u \in T_1} \left(|z - u|^2 - |x - u|^2 \right) \ge \sum_{u \in S_1} \left(|z - u|^2 - |x - u|^2 \right).$$

For $u \in S_1 \cap T_1$, the terms appear on both sides, and we may cancel them, so we are left to prove

$$\sum_{u \in T_1 \setminus S_1} \left(|z - u|^2 - |x - u|^2 \right) \ge \sum_{u \in S_1 \setminus T_1} \left(|z - u|^2 - |x - u|^2 \right)$$

which is true because by the movement of x to z, each term on the left hand side is non negative and each term on the right hand side is non positive.

8.13 Exercises

Exercise 8.1 Construct examples where using distances instead of distance squared gives bad results for Gaussian densities. For example, pick samples from two 1-dimensional unit variance Gaussians, with their centers 10 units apart. Cluster these samples by trial and error into two clusters, first according to k-means and then according to the k-median criteria. The k-means clustering should essentially yield the centers of the Gaussians as cluster centers. What cluster centers do you get when you use the k-median criterion?

Exercise 8.2 Let v = (1,3). What is the L_1 norm of v? The L_2 norm? The square of the L_1 norm?

Exercise 8.3 Show that in 1-dimension, the center of a cluster that minimizes the sum of distances of data points to the center is in general not unique. Suppose we now require the center also to be a data point; then show that it is the median element (not the mean). Further in 1-dimension, show that if the center minimizes the sum of squared distances to the data points, then it is unique.

Exercise 8.4 Construct a block diagonal matrix A with three blocks of size 50. Each matrix element in a block has value p = 0.7 and each matrix element not in a block has value q = 0.3. Generate a 150×150 matrix B of random numbers in the range [0,1]. If $b_{ij} \ge a_{ij}$ replace a_{ij} with the value one. Otherwise replace a_{ij} with value zero. The rows of A have three natural clusters. Generate a random permutation and use it to permute the rows and columns of the matrix A so that the rows and columns of each cluster are randomly distributed.

- 1. Apply the k-mean algorithm to A with k = 3. Do you find the correct clusters?
- 2. Apply the k-means algorithm to A for $1 \le k \le 10$. Plot the value of the sum of squares to the cluster centers versus k. Was three the correct value for k?

Exercise 8.5 Let M be a $k \times k$ matrix whose elements are numbers in the range [0,1]. A matrix entry close to one indicates that the row and column of the entry correspond to closely related items and an entry close to zero indicates unrelated entities. Develop an algorithm to match each row with a closely related column where a column can be matched with only one row.

Exercise 8.6 The simple greedy algorithm of Section 8.3 assumes that we know the clustering radius r. Suppose we do not. Describe how we might arrive at the correct r?

Exercise 8.7 For the k-median problem, show that there is at most a factor of two ratio between the optimal value when we either require all cluster centers to be data points or allow arbitrary points to be centers.

Exercise 8.8 For the k-means problem, show that there is at most a factor of four ratio between the optimal value when we either require all cluster centers to be data points or allow arbitrary points to be centers.

Exercise 8.9 Consider clustering points in the plane according to the k-median criterion, where cluster centers are required to be data points. Enumerate all possible clustering's and select the one with the minimum cost. The number of possible ways of labeling n points, each with a label from $\{1, 2, \ldots, k\}$ is k^n which is prohibitive. Show that we can find the optimal clustering in time at most a constant times $\binom{n}{k} + k^2$. Note that $\binom{n}{k} \leq n^k$ which is much smaller than k^n when $k \ll n$.

Exercise 8.10 Suppose in the previous exercise, we allow any point in space (not necessarily data points) to be cluster centers. Show that the optimal clustering may be found in time at most a constant times n^{2k^2} .

Exercise 8.11 Corollary 8.3 shows that for a set of points $\{a_1, a_2, \ldots, a_n\}$, there is a unique point x, namely their centroid, which minimizes $\sum_{i=1}^{n} |a_i - x|^2$. Show examples where the x minimizing $\sum_{i=1}^{n} |a_i - x|$ is not unique. (Consider just points on the real line.) Show examples where the x defined as above are far apart from each other.

Exercise 8.12 Let $\{\mathbf{a_1}, \mathbf{a_2}, \dots, \mathbf{a_n}\}$ be a set of unit vectors in a cluster. Let $\mathbf{c} = \frac{1}{n} \sum_{i=1}^{n} \mathbf{a_i}$ be the cluster centroid. The centroid \mathbf{c} is not in general a unit vector. Define the similarity between two points \mathbf{a}_i and \mathbf{a}_j as their dot product. Show that the average cluster similarity $\frac{1}{n^2} \sum_{i,j} \mathbf{a_i} \mathbf{a_j}^T$ is the same whether it is computed by averaging all pairs or computing the average similarity of each point with the centroid of the cluster.

Exercise 8.13 For some synthetic data estimate the number of local minima for k-means by using the birthday estimate. Is your estimate an unbaised estimate of the number? an upper bound? a lower bound? Why?

Exercise 8.14 Examine the example in Figure 8.8 and discuss how to fix it. Optimizing according to the k-center or k-median criteria would seem to produce clustering B while clustering A seems more desirable.

Exercise 8.15 Prove that for any two vectors \mathbf{a} and \mathbf{b} , $|\mathbf{a} - \mathbf{b}|^2 \geq \frac{1}{2}|\mathbf{a}|^2 - |\mathbf{b}|^2$.

Exercise 8.16 Let A be an $n \times d$ data matrix, B its best rank k approximation, and C the optimal centers for k-means clustering of rows of A. How is it possible that $||A - B||_F^2 < ||A - C||_F^2$?

Exercise 8.17 Suppose S is a finite set of points in space with centroid $\mu(S)$. If a set T of points is added to S, show that the centroid $\mu(S \cup T)$ of $S \cup T$ is at distance at most $\frac{|T|}{|S|+|T|}|\mu(T) - \mu(S)|$ from $\mu(S)$.

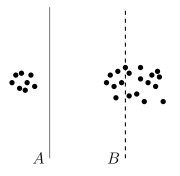


Figure 8.8: insert caption

Exercise 8.18 What happens if we relax this restriction, for example, if we allow for S, the entire set?

Exercise 8.19 Given the graph G = (V, E) of a social network where vertices represent individuals and edges represent relationships of some kind, one would like to define the concept of a community. A number of deferent definitions are possible.

- 1. A subgraph $S = (V_S, E_S)$ whose density $\frac{E_S}{V_s^2}$ is greater than that of the graph $\frac{E}{V^2}$.
- 2. A subgraph S with a low conductance like property such as the number of graph edges leaving the subgraph normalized by the minimum size of S or V S where size is measured by the sum of degrees of vertices in S or in V S.
- 3. A subgraph that has more internal edges than in a random graph with the same degree distribution.

Which would you use and why?

Exercise 8.20 A stochastic matrix is a matrix with non negative entries in which each row sums to one. Show that for a stochastic matrix, the largest eigenvalue is one. Show that the eigenvalue has multiplicity one if and only if the corresponding Markov Chain is connected.

Exercise 8.21 Show that if P is a stochastic matrix and π satisfies $\pi_i p_{ij} = \pi_j p_{ji}$, then for any left eigenvector \mathbf{v} of P, the vector \mathbf{u} with components $u_i = \frac{v_i}{\pi_i}$ is a right eigenvector with the same eigenvalue.

Exercise 8.22 In Theorem (??), how can one clustering $C^{(0)}$ be close to any proper clustering? What if there are several proper clusterings?

Exercise 8.23 Give an example of a clustering problem where the clusters are not linearly separable in the original space, but are separable in a higher dimensional space. **Hint:** Look at the example for Gaussian kernels in the chapter on learning. **Exercise 8.24** The Gaussian kernel maps points to a higher dimensional space. What is this mapping?

Exercise 8.25 Agglomerative clustering requires that one calculate the distances between all pairs of points. If the number of points is a million or more, then this is impractical. One might try speeding up the agglomerative clustering algorithm by maintaining a 100 clusters at each unit of time. Start by randomly selecting a hundred points and place each point in a cluster by itself. Each time a pair of clusters is merged randomly select one of the remaining data points and create a new cluster containing that point. Suggest some other alternatives.

Exercise 8.26 Let A be the adjacency matrix of an undirected graph. Let $d(S, S) = \frac{A(S,S)}{|S|}$ be the density of the subgraph induced by the set of vertices S. Prove that d(S,S) is the average degree of a vertex in S. Recall that $A(S,T) = \sum_{i \in S, j \in T} a_{ij}$

Exercise 8.27 Suppose A is a matrix with non negative entries. Show that A(S,T)/(|S||T|) is maximized by the single edge with highest a_{ij} . Recall that $A(S,T) = \sum_{i \in S, j \in T} a_{ij}$

Exercise 8.28 Suppose A is a matrix with non negative entries and

$$\sigma_1(A) = \mathbf{x}^T A \mathbf{y} = \sum_{i,j} x_i a_{ij} y_j, \qquad |\mathbf{x}| = |\mathbf{y}| = 1.$$

Zero out all x_i less than $1/2\sqrt{n}$ and all y_j less than $1/2\sqrt{d}$. Show that the loss is no more than $1/4^{th}$ of $\sigma_1(A)$.

Exercise 8.29 Consider other measures of density such as $\frac{A(S,T)}{|S|^{\rho}|T|^{\rho}}$ for different values of ρ . Discuss the significance of the densest subgraph according to these measures.

Exercise 8.30 Let A be the adjacency matrix of an undirected graph. Let M be the matrix whose ij^{th} element is $a_{ij} - \frac{d_i d_j}{2m}$. Partition the vertices into two groups S and \bar{S} . Let s be the indicator vector for the set S and let \bar{s} be the indicator variable for \bar{S} . Then $s^T M s$ is the number of edges in S above the expected number given the degree distribution and $s^T M \bar{s}$ is the number of edges from S to \bar{S} above the expected number given the degree distribution. Prove that if $s^T M s$ is positive $s^T M \bar{s}$ must be negative.

Exercise 8.31 Which of the three axioms, scale invariance, richness, and consistency are satisfied by the following clustering algorithms.

- 1. k-means
- 2. Spectral Clustering.

Exercise 8.32 (*Research Problem*): What are good measures of density that are also effectively computable? Is there empirical/theoretical evidence that some are better than others?

9 Topic Models, Hidden Markov Process, Graphical Models, and Belief Propagation

In the chapter on learning and VC dimension, we saw many model-fitting problems. There we were given labeled data and simple classes of functions: half-spaces, support vector machines, etc. The problem was to fit the best model from a class of functions to the data. Model fitting is of course more general and in this chapter we discuss some useful models. These general models are often computationally infeasible, in the sense that they do not admit provably efficient algorithms. Nevertheless, data often falls into special cases of these models that can be solved efficiently.

9.1 Topic Models

A *topic model* is a model for representing a large collection of documents. Each document is viewed as a combination of topics and each topic has a set of word frequencies. For a collection of news articles over a period, the topics may be politics, sports, science, etc. For the topic politics, the words like "president" and "election" may have high frequencies and for the topic sports, words like "batter" and "goal" may have high frequencies. A news item document may be 60% on politics and 40% on sports. The word frequencies in the document will be a convex combination of word frequencies for the two topics, politics and sports, with weights 0.6 and 0.4 respectively. We describe this more formally with vectors and matrices.

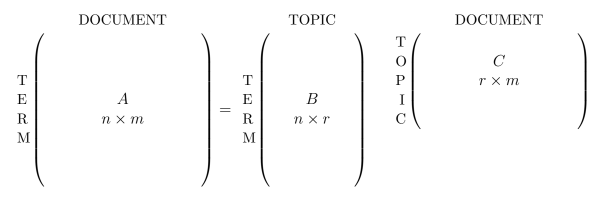
Each document is viewed as a "bag of words". We disregard the order and context in which each term occurs in the document and instead only list the frequency of occurrences of each term. Frequency is the number of occurrences of the term divided by the total number of all terms in the document. Discarding context information may seem wasteful, but this approach works well in practice and is widely used. Each document is an *n*-dimensional vector where n is the total number of different terms in all the documents in the collection. Each component of the vector is the frequency of a particular term in the document. Terms are words or phrases. Not all words are chosen as terms; articles, simple verbs, and pronouns like "a", "is", and "it" may be ignored. /newline

Represent the collection of documents by a $n \times m$ matrix A, called the *term-document* matrix, with one column per document in the collection. The topic model hypothesizes that there are r topics and each of the m documents is a combination of topics. The number of topics r is usually much smaller than the number of terms n. So corresponding to each document, there is a vector with r components telling us the fraction of the document that is on each of the topics. In the example above, this vector will have 0.6 in the component for politics and 0.4 in the component for sports. Arrange these vectors as the columns of a $r \times m$ matrix C, called the *topic-document* matrix. There is a third matrix B which is $n \times r$. Each column of B corresponds to a topic; each component of the column gives the frequency of a term in that topic. In the simplest model, the term frequencies in documents are exact combinations of term frequencies in the various topics

that make up the document. So, a_{ij} , the frequency of the i^{th} term in the j^{th} document is the sum over all topics l of the fraction of document j which is on topic l times the frequency of term i in topic l. In matrix notation,

$$A = BC.$$

Pictorially, we can represent this as:



This model is too simple to be realistic since the frequency of each term in the document is unlikely to be exactly what is given by the equation A = BC. So, a more sophisticated stochastic model is used in practice.

From the document collection we observe the $n \times m$ matrix A. Can we find B and C such that A = BC? The top r singular vectors from a singular value decomposition of A give a factorization BC. But there are additional constraints stemming from the fact that frequencies of terms in one particular topic are nonnegative reals summing to one and from the fact that the fraction of each topic a particular document is on are also nonnegative reals summing to one. Altogether the constraints are:

- 1. A = BC and $\sum_{i} a_{ij} = 1$.
- 2. The entries of B and C are all non-negative.
- 3. Each column of B and each column of C sums to one.

Given the first two conditions, we can achieve the third by multiplying the i^{th} column of B by a positive real number and dividing the i^{th} row of C by the same real number without violating A = BC. By doing this, one may assume that each column of B sums to one. Since $\sum_i a_{ij}$ is the total frequency of all terms in document j, $\sum_i a_{ij} = 1$. Now $a_{ij} = \sum_k b_{ik}c_{kj}$ implies $\sum_i a_{ij} = \sum_{i,k} b_{ik}c_{kj} = \sum_k c_{kj} = 1$. Thus, the columns of C also sum to one.

The problem can be posed as one of factoring the given matrix A into the product of two matrices with nonnegative entries called nonnegative matrix factorization.

Nonnegative matrix factorization (NMF) Given an $n \times m$ matrix A and an integer r, determine whether there is a factorization of A into XY where, X is an $n \times r$ matrix with nonnegative entries and Y is $r \times m$ matrix with nonnegative entries and if so, find such a factorization.

Nonnegative matrix factorization is a more general problem than topic modeling and there are many heuristic algorithms to solve the problem. But in general, they suffer from one of two problems, they can get stuck at local optima that are not solutions or take exponential time. In fact, the general NMF problem is NP-hard. In practice, often r is much smaller than n and m. We first show that while the NMF problem as formulated above is a nonlinear problem in r(n + m) unknown entries of X and Y, it can be reformulated as a nonlinear problem with just $2r^2$ unknowns under the simple nondegeneracy assumption that A has rank r. Think of r as say, 25, while n and m are in the tens of thousands to see why this is useful.

Lemma 9.1 If A has rank r, then the NMF problem can be formulated as a problem with $2r^2$ unknowns.

Proof: If A = XY, then each row of A is a linear combination of the rows of Y. So the space spanned by the rows of A must be contained in the space spanned by the rows of Y. The latter space has dimension at most r, while the former has dimension exactly r. So they must be equal. Thus, every row of Y must be a linear combination of the rows of A. Choose any set of r independent rows of A to form a $r \times m$ matrix A_1 . Then $Y = SA_1$ for some $r \times r$ matrix S. By analogous reasoning, if A_2 is a $n \times r$ matrix of r independent columns of A, there is a $r \times r$ matrix T such that $X = A_2T$. Now we can easily cast NMF in terms of the unknowns S and T.

$$A = A_2 T S A_1 \qquad (SA_1)_{ij} \ge 0 \qquad (A_2 T)_{kl} \ge 0 \qquad \forall i, j, k, l.$$

It remains to solve the nonlinear problem in the $2r^2$ variables. There is a classical algorithm that solves such problems in time exponential in r^2 and polynomial in the other parameters. In fact, there is a logical theory, called the theory of reals of which this is a special case and any problem in the theory of reals can be solved in time exponential only in the number of variables. We do not give details here.

Besides the special case when r is small, there is another important case of NMF in the topic modeling application that can be solved. This is the case when there are *anchor terms*. An anchor term for a topic is a term that occurs in the topic and does not occur in any other topic. For example, the term "batter" may an anchor term for the topic baseball and "election" for the topic politics. Consider the case when each topic has an anchor term. In matrix notation, this assumes that for each column of the term-topic matrix B, there is a row whose sole nonzero entry is in that column. In this case, it is easy to see that each row of the topic-document matrix C has a scaled copy of it occurring as a row of the given term-document matrix A. Here is an illustrative diagram:

$$\begin{pmatrix} 0.3c_4 \\ 0.2c_2 \end{pmatrix} = \begin{array}{c} \text{election} \begin{pmatrix} 0 & 0 & 0 & 0.3 \\ 0 & 0.2 & 0 & 0 \\ 0 & 0.2 & 0 & 0 \\ A \end{pmatrix} \begin{pmatrix} c_1 \\ c_2 \\ c_3 \\ \vdots \\ c_n \end{pmatrix}$$

If we knew which rows of A were copies of rows of C, called special rows of A, we could find C. Once C is known, we could solve the linear equations A = BC and inequalities $b_{ij} \ge 0$, to get B. The following lemma shows that after making one modification, we can find the rows of A that are special. Suppose a row of C is a nonnegative linear combination of the other rows of C. Eliminate that row of C as well as the corresponding column of B, suitably modifying the other columns of B, maintaining A = BC. For example, if row 5 of C equals 4 times row 3 of C plus 3 times row 6 of C, then we can delete row 5 of C, then add 4 times column 5 of B to column 3 of B and add 3 times column 5 of B

Repeating this until each row of C is positively independent of the other rows of C, i.e., it cannot be expressed as a nonnegative linear combination of the other rows. We still have a scaled copy of each row of C in A. Further, the other rows of A are all nonnegative linear combinations of rows of C and thus are nonnegative linear combinations of the special rows of A.

Lemma 9.2 Suppose A has a factorization A = BC, where the rows of C are positively independent and for each column of B, there is a row that has its sole nonzero entry in that column. Then there is a scaled copy of each row of C in A and furthermore, the rows of A that are scaled copies of rows of C are precisely the rows of A that are positively independent of other rows of A. These rows can be identified by solving a linear program, one program per row.

Proof: The set of special rows of A can be identified by solving n linear programming problems. Check each row of A to see if it is positively independent of all other rows. Denote by \mathbf{a}_i the i th row of A. Then, the i^{th} row is positively dependent upon the others if and only if there are real numbers $x_1, x_2, \ldots x_{i-1}, x_{i+1}, \ldots x_n$ such that

$$\sum_{j \neq i} x_j \mathbf{a_j} = \mathbf{a_i}, \quad x_j \ge 0$$

This is a linear program.

As we remarked earlier, the equation A = BC will not hold exactly. A more practical model views A as a matrix of probabilities rather than exact frequencies. In this model, each document is generated by picking its terms in independent trials. Each trial for document j picks term 1 with probability a_{1j} ; term 2 with probability a_{2j} , etc. We are not given entire documents; instead we are given s independent trials for each document. Our job is to find B and C. We do not discuss the details of either the model or the algorithms. In this new situation, algorithms are known to find B and C when there exist anchor terms, even with a small number s of trials.

At the heart of such an algorithm is the following problem:

Approximate NMF Given a $n \times m$ matrix A and the promise that there is a $n \times r$ matrix B and a $r \times m$ matrix C, both with nonnegative entries, such that $||A - BC||_F \leq \Delta$, find B' and C' of the same dimensions, with nonnegative entries such that $||A - B'C||_F \leq \Delta'$.

Here, Δ' is related to Δ and if the promise does not hold, the algorithm is allowed to return any answer.

Now for the case when anchor words exist, this reduces to the problem of finding which rows of A have the property that no point close to the row is positively dependent on other rows. It is easy to write the statement that there is a vector \mathbf{y} close to \mathbf{a}_i which is positively dependent on the other rows as a convex program:

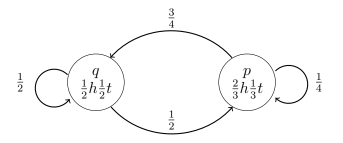
$$\exists x_1, x_2, \dots, x_{i-1}, x_{i+1}, \dots, x_n \text{ such that } \left| \sum_{j \neq i} x_j \mathbf{a_j} - \mathbf{a_i} \right| \leq \varepsilon$$

 $|\sum_{j\neq i} x_j \mathbf{a}_j - \mathbf{a}_i|$ is convex function of x_j and hence this problem can be solved efficiently.

9.2 Hidden Markov Model

A hidden Markov model, HMM, consists of a finite set of states with a transition between each pair of states. There is an initial probability distribution α on the states and a transition probability a_{ij} associated with the transition from state *i* to state *j*. Each state has a probability distribution p(O, i) giving the probability of outputting the symbol O in state *i*. A transition consists of two components. A state transition to a new state followed by the output of a symbol. The HMM starts by selecting a start state according to the distribution α and outputting a symbol.

Example: An example of a HMM is the graph with two states q and p illustrated below.



The initial distribution is $\alpha(q) = 1$ and $\alpha(p) = 0$. At each step a change of state occurs followed by the output of heads or tails with probability determined by the new state.

We consider three problems in increasing order of difficulty. First, given a HMM what is the probability of a given output sequence? Second, given a HMM and an output sequence, what is the most likely sequence of states? And third, knowing that the HMM has at most n states and given an output sequence, what is the most likely HMM? Only the third problem concerns a "hidden" Markov model. In the other two problems, the model is known and the questions can be answered in polynomial time using dynamic programming. There is no known polynomial time algorithm for the third question.

How probable is an output sequence

Given a HMM, how probable is the output sequence $O = O_0O_1O_2\cdots O_T$ of length T+1? To determine this, calculate for each state i and each initial segment of the sequence of observations, $O_0O_1O_2\cdots O_t$ of length t+1, the probability of observing $O_0O_1O_2\cdots O_t$ ending in state i. This is done by a dynamic programming algorithm starting with t = 0 and increasing t. For t = 0 there have been no transitions. Thus, the probability of observing O_0 ending in state i is the initial probability of observing $O_0O_1O_2\cdots O_t$ ending in state i is the sum of the probabilities over all states j of observing $O_0O_1O_2\cdots O_{t-1}$ ending in state j times the probability of going from state j to state i and observing O_t . The time to compute the probability of a sequence of length T when there are n states is $O(n^2T)$. The factor n^2 comes from the calculation for each time unit of the contribution from each possible previous state to the probability of each possible current state. The space complexity is O(n) since one only needs to remember the probability of reaching each state for the most recent value of t.

Algorithm to calculate the probability of the output sequence

The probability, $\operatorname{Prob}(O_0O_1\cdots O_T, i)$ of the output sequence $O_0O_1\cdots O_T$ ending in state *i* is given by

 $Prob(O_0, i) = \alpha(i)p(O_0, i)$

for t = 1 to T

$$\operatorname{Prob}(O_0O_1\cdots O_t, i) = \sum_j \operatorname{Prob}(O_0O_1\cdots O_{t-1}, j)a_{ij}p(O_{t+1}, i)$$

Example: What is the probability of the sequence hhht by the HMM in the above two state example?

t = 3	$\frac{3}{32}\frac{1}{2}\frac{1}{2} + \frac{5}{72}\frac{3}{4}\frac{1}{2} = \frac{19}{384}$	$\frac{3}{32}\frac{1}{2}\frac{1}{3} + \frac{5}{72}\frac{1}{4}\frac{1}{3} = \frac{37}{64\times27}$
t=2	$\frac{1}{8}\frac{1}{2}\frac{1}{2} + \frac{1}{6}\frac{3}{4}\frac{1}{2} = \frac{3}{32}$	$\frac{1}{8}\frac{1}{2}\frac{2}{3} + \frac{1}{6}\frac{1}{4}\frac{2}{3} = \frac{5}{72}$
t = 1	$\frac{1}{2}\frac{1}{2}\frac{1}{2} = \frac{1}{8}$	$\frac{1}{2}\frac{1}{2}\frac{2}{3} = \frac{1}{6}$
t = 0	$\frac{1}{2}$	0
	q	p

For t = 0, the q entry is $\frac{1}{2}$ since the probability of being in state q is one and the probability of outputting heads is $\frac{1}{2}$. The entry for p is zero since the probability of starting in state p is zero. For t = 1, the q entry is $\frac{1}{8}$ since for t = 0 the q entry is $\frac{1}{2}$ and in state q the HMM goes to state q with probability $\frac{1}{2}$ and outputs heads with probability $\frac{1}{2}$. The p entry is $\frac{1}{6}$ since for t = 0 the q entry is $\frac{1}{2}$ and in state p with probability $\frac{1}{2}$ and outputs heads with probability $\frac{1}{2}$. The p entry is $\frac{1}{6}$ since for t = 0 the q entry is $\frac{1}{2}$ and in state q the HMM goes to state p with probability $\frac{1}{2}$ and outputs heads with probability $\frac{2}{3}$. For t = 2, the q entry is $\frac{3}{32}$ which consists of two terms. The first term is the probability of ending in state q at t = 1 times the probability of staying in q and outputting h. The second is the probability of ending in state p at t = 1 times the probability of going from state p to state q and outputting h.

From the table, the probability of producing the sequence hhht is $\frac{19}{384} + \frac{37}{1728} = 0.0709$.

The most likely sequence of states - the Viterbi algorithm

Given a HMM and an observation $O = O_0O_1 \cdots O_T$, what is the most likely sequence of states? The solution is given by the Viterbi algorithm, which is a slight modification to the dynamic programming algorithm just given for determining the probability of an output sequence. For $t = 0, 1, 2, \ldots, T$ and for each state *i*, calculate the probability of the most likely sequence of states to produce the output $O_0O_1O_2\cdots O_t$ ending in state *i*. For each value of *t*, calculate the most likely sequence of states by selecting over all states *j* the most likely sequence producing $O_0O_1O_2\cdots O_t$ and ending in state *i* consisting of the most likely sequence producing $O_0O_1O_2\cdots O_t$ and ending in state *j* followed by the transition from *j* to *i* producing O_t . Note that in the previous example, we added the probabilities of each possibility together. Now we take the maximum and also record where the maximum came from. The time complexity is $O(n^2T)$ and the space complexity is O(nT). The space complexity bound is argued as follows. In calculating the probability of the most likely sequence of states that produces $O_0O_1 \ldots O_t$ ending in state *i*, we remember the previous state j by putting an arrow with edge label t from i to j. At the end, can find the most likely sequence by tracing backwards as is standard for dynamic programming algorithms.

Example: For the earlier example what is the most likely sequence of states to produce the output hhht?

t = 3	$\max\{\frac{1}{48}\frac{1}{2}\frac{1}{2}, \frac{1}{24}\frac{3}{4}\frac{1}{2}\} = \frac{1}{64} q \text{ or } p$	$\max\{\frac{3}{48}\frac{1}{2}\frac{1}{3}, \frac{1}{24}\frac{1}{4}\frac{1}{3}\} = \frac{1}{96} q$
t = 2	$\max\{\frac{1}{8}\frac{1}{2}\frac{1}{2}, \frac{1}{6}\frac{3}{4}\frac{1}{2}\} = \frac{3}{48} p$	$\max\{\frac{1}{8}\frac{1}{2}\frac{2}{3}, \frac{1}{6}\frac{1}{4}\frac{2}{3}\} = \frac{1}{24} q$
t = 1	$\frac{1}{2}\frac{1}{2}\frac{1}{2} = \frac{1}{8}$ q	$\frac{1}{2}\frac{1}{2}\frac{2}{3} = \frac{1}{6}$ q
t = 0	$\frac{1}{2}$ q	0 р
	q	p

Note that the two sequences of states, qqpq and qpqq, are tied for the most likely sequences of states.

Determining the underlying hidden Markov model

Given an *n*-state HMM, how do we adjust the transition probabilities and output probabilities to maximize the probability of an output sequence $O_1O_2\cdots O_T$? The assumption is that T is much larger than n. There is no known computationally efficient method for solving this problem. However, there are iterative techniques that converge to a local optimum.

Let a_{ij} be the transition probability from state *i* to state *j* and let $b_j(O_k)$ be the probability of output O_k given that the HMM is in state *j*. Given estimates for the HMM parameters, a_{ij} and b_j , and the output sequence O, we can improve the estimates by calculating for each unit of time the probability that the HMM goes from state *i* to state *j* and outputs the symbol O_k .

Given estimates for the HMM parameters, a_{ij} and b_j , and the output sequence O, the probability $\delta_t(i, j)$ of going from state *i* to state *j* at time *t* is given by the probability of producing the output sequence O and going from state *i* to state *j* at time *t* divided by the probability of producing the output sequence O.

$$\delta_t(i,j) = \frac{a_t(i)a_{ij}b_j(O_{t+1})\beta_{t+1}(j)}{p(O)}$$

The probability p(O) is the sum over all pairs of states *i* and *j* of the numerator in the above formula for $\delta_t(i, j)$. That is,

$$p(O) = \sum_{i} \sum_{j} \alpha_{t}(j) a_{ij} b_{j}(O_{t+1}) \beta_{t+1}(j).$$

a_{ij}	transition probability from state i to state j
$b_j(O_{t+1})$	probability of O_{t+1} given that the HMM is in state j at time $t+1$
$\alpha_t(i)$	probability of seeing $O_0 O_1 \cdots O_t$ and ending in state <i>i</i> at time <i>t</i>
$\beta_{t+1}(j)$	probability of seeing the tail of the sequence $O_{t+2}O_{t+3}\cdots O_T$ given state j at time $t+1$
$\delta(i,j)$	probability of going from state i to state j at time t given the sequence of outputs O
$s_t(i)$	probability of being in state i at time t given the sequence of outputs O
p(O)	probability of output sequence O

The probability of being in state i at time t is given by

$$s_t(i) = \sum_{j=1}^n \delta_t(i,j).$$

Note that $\delta_t(i, j)$ is the probability of being in state *i* at time *t* given $O_0O_1O_2\cdots O_t$ but it is not the probability of being in state *i* at time *t* given *O* since it does not take into account the remainder of the sequence *O*. Summing $s_t(i)$ over all time periods gives the expected number of times state *i* is visited and the sum of $\delta_t(i, j)$ over all time periods gives the expected number of times edge *i* to *j* is traversed.

Given estimates of the HMM parameters $a_{i,j}$ and $b_j(O_k)$, we can calculate by the above formulas estimates for

- 1. $\sum_{i=1}^{T-1} s_i(i)$, the expected number of times state *i* is visited and departed from
- 2. $\sum_{i=1}^{T-1} \delta_t(i,j)$, the expected number of transitions from state *i* to state *j*

Using these estimates we can obtain new estimates of the HMM parameters

$$\overline{a_{ij}} = \frac{\text{expected number of transitions from state } i \text{ to state } j}{\text{expected number of transitions out of state } i} = \frac{\sum_{t=1}^{T-1} \delta_t(i,j)}{\sum_{t=1}^{T-1} s_t(i)}$$
$$\overline{b_j}(O_k) = \frac{\text{expected number of times in state } j \text{ observing symbol } O_k}{\text{expected number of times in state } j} = \frac{O_k}{\sum_{t=1}^{T-1} s_t(j)}$$

By iterating the above formulas we can arrive at a local optimum for the HMM parameters $a_{i,j}$ and $b_j(O_k)$.

9.3 Graphical Models, and Belief Propagation

A graphical model is a compact representation of a function of n variables x_1, x_2, \ldots, x_n . It consists of a graph, directed or undirected, whose vertices correspond to variables that take on values from some set. In this chapter, we consider the case where the function is a probability distribution and the set of values the variables take on is finite, although graphical models are often used to represent probability distributions with continuous variables. The edges of the graph represent relationships or constraints between the variables.

The directed model represents a joint probability distribution that factors into a product of conditional probabilities.

$$p(x_1, x_2, \dots, x_n) = \prod_{i=1}^n p(x_i | \text{parents of } x_i)$$

It is assumed that the directed graph is acyclic. The directed graphical model is called a *Bayesian* or *belief network* and appears frequently in the artificial intelligence and the statistics literature.

The undirected graph model, called a Markov random field, can also represent a joint probability distribution of the random variables at its vertices. In many applications the Markov random field represents a function of the variables at the vertices which is to be optimized by choosing values for the variables.

A third model called the factor model is akin to the Markov random field, but here the dependency sets have a different structure. In the following sections we describe all these models in more detail.

9.4 Bayesian or Belief Networks

A Bayesian network is a directed acyclic graph where vertices correspond to variables and a directed edge from y to x represents a conditional probability p(x|y). If a vertex x has edges into it from y_1, y_2, \ldots, y_k , then the conditional probability is $p(x | y_1, y_2, \ldots, y_k)$. The variable at a vertex with no in edges has an unconditional probability distribution. If the value of a variable at some vertex is known, then the variable is called evidence. An important property of a Bayesian network is that the joint probability is given by the product over all nodes of the conditional probability of the node conditioned on all its immediate predecessors.

In the example of Fig. 9.1, a patient is ill and sees a doctor. The doctor ascertains the symptoms of the patient and the possible causes such as whether the patient was in

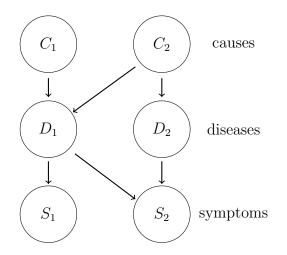


Figure 9.1: A Bayesian network

contact with farm animals, whether he had eaten certain foods, or whether the patient has an hereditary predisposition to any diseases. Using the above Bayesian network where the variables are true or false, the doctor may wish to determine one of two things. What is the marginal probability of a given disease or what is the most likely set of diseases. In determining the most likely set of diseases, we are given a T or F assignment to the causes and symptoms and ask what assignment of T or F to the diseases maximizes the joint probability. This latter problem is called the maximum a posteriori probability (MAP).

Given the conditional probabilities and the probabilities $p(C_1)$ and $p(C_2)$ in Figure 9.1, the joint probability $p(C_1, C_2, D_1, ...)$ can be computed easily for any combination of values of $C_1, C_2, D_1, ...$ However, we might wish to find that value of the variables of highest probability (MAP) or we might want one of the marginal probabilities $p(D_1)$ or $p(D_2)$. The obvious algorithms for these two problems require evaluating the probability $p(C_1, C_2, D_1, ...)$ over exponentially many input values or summing the probability $p(C_1, C_2, D_1, ...)$ over exponentially many values of the variables other than those for which we want the marginal probability. In certain situations, when the joint probability distribution can be expressed as a product of factors, a belief propagation algorithm can solve the maximum a posteriori problem or compute all marginal probabilities quickly.

9.5 Markov Random Fields

The Markov random field model arose first in statistical mechanics where it was called the Ising model. It is instructive to start with a description of it. The simplest version of the Ising model consists of n particles arranged in a rectangular $\sqrt{n} \times \sqrt{n}$ grid. Each particle can have a spin that is denoted ± 1 . The energy of the whole system depends on interactions between pairs of neighboring particles. Let x_i be the spin, ± 1 , of the i^{th} particle. Denote by $i \sim j$ the relation that i and j are adjacent in the grid. In the Ising model, the energy of the system is given by

$$f(x_1, x_2, \dots, x_n) = \exp\left(c\sum_{i\sim j} |x_i - x_j|\right).$$

The constant c can be positive or negative. If c < 0, then energy is lower if many adjacent pairs have opposite spins and if c > 0 the reverse holds. The model was first used to model probabilities of spin configurations. The hypothesis was that for each $\{x_1, x_2, \ldots, x_n\}$ in $\{-1, +1\}^n$, the energy of the configuration with these spins is proportional to $f(x_1, x_2, \ldots, x_n)$.

In most computer science settings, such functions are mainly used as objective functions that are to be optimized subject to some constraints. The problem is to find the minimum energy set of spins under some constraints on the spins. Usually the constraints just specify the spins of some particles. Note that when c > 0, this is the problem of minimizing $\sum_{i\sim j} |x_i - x_j|$ subject to the constraints. The objective function is convex and so this can be done efficiently. If c < 0, however, we need to minimize a concave function for which there is no known efficient algorithm. The minimization of a concave function in general is NP-hard.

A second important motivation comes from the area of vision. It has to to do with reconstructing images. Suppose we are given observations of the intensity of light at individual pixels, x_1, x_2, \ldots, x_n , and wish to compute the true values, the true intensities, of these variables y_1, y_2, \ldots, y_n . There may be two sets of constraints, the first stipulating that the y_i must be close to the corresponding x_i and the second, a term correcting possible observation errors, stipulating that y_i must be close to the values of y_j for $j \sim i$. This can be formulated as

$$\min_{\mathbf{y}} \left(\sum_{i} |x_i - y_i| + \sum_{i \sim j} |y_i - y_j| \right),$$

where the values of x_i are constrained to be the observed values. The objective function is convex and polynomial time minimization algorithms exist. Other objective functions using say sum of squares instead of sum of absolute values can be used and there are polynomial time algorithms as long as the function to be minimized is convex.

More generally, the correction term may depend on all grid points within distance two of each point rather than just immediate neighbors. Even more generally, we may have nvariables $y_1, y_2, \ldots y_n$ with the value of some already specified and subsets $S_1, S_2, \ldots S_m$ of these variables constrained in some way. The constraints are accumulated into one objective function which is a product of functions f_1, f_2, \ldots, f_m , where function f_i is evaluated on the variables in subset S_i . The problem is to minimize $\prod_{i=1}^m f_i(y_j, j \in S_i)$ subject to constrained values. Note that the vision example had a sum instead of a product, but by taking exponentials we can turn the sum into a product as in the Ising model.

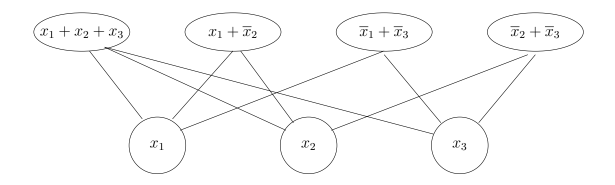


Figure 9.2: The factor graph for the function $f(x_1, x_2, x_3) = (x_1 + x_2 + x_3)(x_1 + \bar{x}_2)(x_1 + \bar{x}_3)(\bar{x}_2 + \bar{x}_3).$

In general, the f_i are not convex; indeed they may be discrete. So the minimization cannot be carried out by a known polynomial time algorithm. The most used forms of the Markov random field involve S_i which are cliques of a graph. So we make the following definition.

A *Markov Random Field* consists of an undirected graph and an associated function that factorizes into functions associated with the cliques of the graph. The special case when all the factors correspond to cliques of size one or two is of interest.

9.6 Factor Graphs

Factor graphs arise when we have a function f of a variables $\mathbf{x} = (x_1, x_2, \ldots, x_n)$ that can be expressed as $f(\mathbf{x}) = \prod_{\alpha} f_{\alpha}(\mathbf{x}_{\alpha})$ where each factor depends only on some small number of variables \mathbf{x}_{α} . The difference from Markov random fields is that the variables corresponding to factors do not necessarily form a clique. Associate a bipartite graph where one set of vertices correspond to the factors and the other set to the variables. Place an edge between a variable and a factor if the factor contains that variable. See Figure 9.2

9.7 Tree Algorithms

Let $f(\mathbf{x})$ be a function that is a product of factors. When the factor graph is a tree there are efficient algorithms for solving certain problems. With slight modifications, the algorithms presented can also solve problems where the function is the sum of terms rather than a product of factors.

The first problem is called *marginalization* and involves evaluating the sum of f over all variables except one. In the case where f is a probability distribution the algorithm

computes the marginal probabilities and thus the word marginalization. The second problem involves computing the assignment to the variables that maximizes the function f. When f is a probability distribution, this problem is the maximum a posteriori probability or MAP problem.

If the factor graph is a tree, then there exists an efficient algorithm for solving these problems. Note that there are four problems: the function f is either a product or a sum and we are either marginalizing or finding the maximizing assignment to the variables. All four problems are solved by essentially the same algorithm and we present the algorithm for the marginalization problem when f is a product. Assume we want to "sum out" all the variables except x_1 , leaving a function of x_1 .

Call the variable node associated with the variable x_i node x_i . First, make the node x_1 the root of the tree. It will be useful to think of the algorithm first as a recursive algorithm and then unravel the recursion. We want to compute the product of all factors occurring in the sub-tree rooted at the root with all variables except the root-variable summed out. Let g_i be the product of all factors occurring in the sub-tree rooted at node x_i with all variables occurring in the subtree except x_i summed out. Since this is a tree, x_1 will not reoccur anywhere except the root. Now, the grandchildren of the root are variable nodes and suppose for recursion, each grandchild x_i of the root, has already computed its g_i . It is easy to see that we can compute g_1 as follows.

Each grandchild x_i of the root passes its g_i to its parent, which is a factor node. Each child of x_1 collects all its children's g_i , multiplies them together with its own factor and sends the product to the root. The root multiplies all the products it gets from its children and sums out all variables except its own variable, namely here x_1 .

Unraveling the recursion is also simple, with the convention that a leaf node just receives 1, product of an empty set of factors, from its children. Each node waits until it receives a message from each of its children. After that, if the node is a variable node, it computes the product of all incoming messages, and sums this product function over all assignments to the variables except for the variable of the node. Then, it sends the resulting function of one variable out along the edge to its parent. If the node is a factor node, it computes the product of its factor function along with incoming messages from all the children and sends the resulting function out along the edge to its parent.

The reader should prove that the following invariant holds assuming the graph is a tree:

Invariant The message passed by each variable node to its parent is the product of all factors in the subtree under the node with all variables in the subtree except its own summed out.

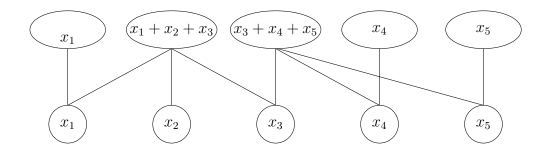


Figure 9.3: The factor graph for the function $f = x_1 (x_1 + x_2 + x_3) (x_3 + x_4 + x_5) x_4 x_5$.

Consider the following example where

$$f = x_1 \left(x_1 + x_2 + x_3 \right) \left(x_3 + x_4 + x_5 \right) x_4 x_5$$

and the variables take on values 0 or 1. Consider marginalizing f by computing

$$f(x_1) = \sum_{x_2 x_3 x_4 x_5} x_1 (x_1 + x_2 + x_3) (x_3 + x_4 + x_5) x_4 x_5,$$

In this case the factor graph is a tree as shown in Figure 9.3. The factor graph as a rooted tree and the messages passed by each node to its parent are shown in Figure 9.4. If instead of computing marginal's, one wanted the variable assignment that maximizes the function f, one would modify the above procedure by replacing the summation by a maximization operation. Obvious modifications handle the situation where $f(\mathbf{x})$ is a sum of products.

$$f\left(\mathbf{x}\right) = \sum_{x_1,\dots,x_n} g\left(\mathbf{x}\right)$$

9.8 Message Passing in general Graphs

The simple message passing algorithm in the last section gives us the one variable function of x_1 when we sum out all the other variables. For a general graph that is not a tree, we formulate an extension of that algorithm. But unlike the case of trees, there is no proof that the algorithm will converge and even if it does, there is no guarantee that the limit is the marginal probability. This has not prevented its usefulness in some applications.

First, lets ask a more general question, just for trees. Suppose we want to compute for each *i* the one variable function of x_i when we sum out all variables $x_j, j \neq i$. Do we have to repeat what we did for x_1 once for each x_i ? Luckily, the answer is no. It will suffice to do a second pass from the root to the leaves of essentially the same message passing algorithm to get all the answers. Recall that in the first pass, each edge of the tree has sent a message "up", from the child to the parent. In the second pass, each edge will send a message down from the parent to the child. We start with the root and work

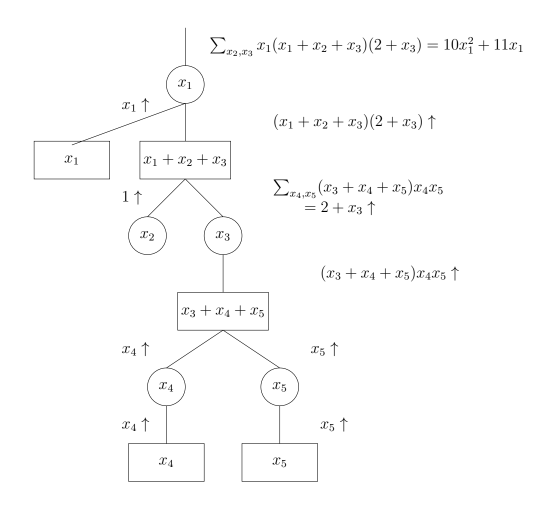


Figure 9.4: Messages.

downwards for this pass. Each node waits until its parent has sent it a message before sending messages to each of its children. The rules for messages are:

Rule 1 The message from a factor node v to a child x_i , which is the variable node x_i , is the product of all messages received by v in both passes from all nodes other than x_i times the factor at v itself.

Rule 2 The message from a variable node x_i to a factor node child, v, is the product of all messages received by x_i in both passes from all nodes except v, with all variables except x_i summed out. The message is a function of x_i alone.

At termination, when the graph is a tree, if we take the product of all messages received in both passes by a variable node x_i and sum out all variables except x_i in this product, what we get is precisely the entire function marginalized to x_i . We do not give the proof here. But the idea is simple. We know from the first pass that the product of the messages coming to a variable node x_i from its children is the product of all factors in the sub-tree rooted at x_i . In the second pass, we claim that the message from the parent v to x_i is the product of all factors which are not in the sub-tree rooted at x_i which one can show either directly or by induction working from the root downwards.

We can apply the same rules 1 and 2 to any general graph. We do not have child and parent relationships and it is not possible to have the two synchronous passes as before. The messages keep flowing and one hopes that after some time, the messages will stabilize, but nothing like that is proven. We state the algorithm for general graphs now:

Rule 1 At each unit of time, each factor node v sends a message to each adjacent node x_i . The message is the product of all messages received by v at the previous step except for the one from x_i multiplied by the factor at v itself.

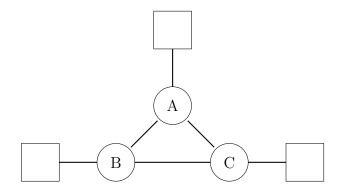
Rule 2 At each time, each variable node x_i sends a message to each adjacent node v. The message is the product of all messages received by x_i at the previous step except the one from v, with all variables except x_i summed out.

9.9 Graphs with a Single Cycle

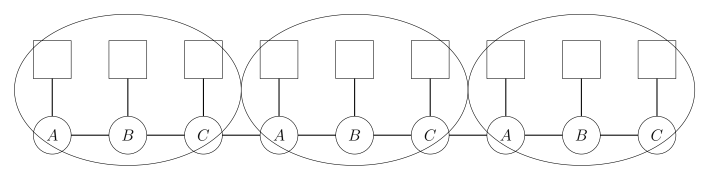
The message passing algorithm gives the correct answers on trees and on certain other graphs. One such situation is graphs with a single cycle which we treat here. We switch from the marginalization problem to the MAP problem as the proof of correctness is simpler for the MAP problem. Consider the network in the Figure 9.5a with a single cycle. The message passing scheme will multiply count some evidence. The local evidence at A will get passed around the loop and will come back to A. Thus, A will count the local evidence multiple times. If all evidence is multiply counted in equal amounts, then there is a possibility that all though the numerical values of the marginal probabilities (beliefs) are wrong, the algorithm still converges to the correct maximum a posteriori assignment.

Consider the unwrapped version of the graph in Figure 9.5b. The messages that the loopy version will eventually converge to, assuming convergence, are the same messages that occur in the unwrapped version provided that the nodes are sufficiently far in from the ends. The beliefs in the unwrapped version are correct for the unwrapped graph since it is a tree. The only question is, how similar are they to the true beliefs in the original network.

Write $p(A, B, C) = e^{\log p(A, B, C)} = e^{J(A, B, C)}$ where $J(A, B, C) = \log p(A, B, C)$. Then the probability for the unwrapped network is of the form $e^{kJ(A, B, C)+J'}$ where the J' is associated with vertices at the ends of the network where the beliefs have not yet stabilized and the kJ(A, B, C) comes from k inner copies of the cycle where the beliefs have stabilized. Note that the last copy of J in the unwrapped network shares an edge with J'and that edge has an associated Ψ . Thus, changing a variable in J has an impact on the



(a) A graph with a single cycle



(b) Segment of unrolled graph

Figure 9.5: Unwrapping a graph with a single cycle

value of J' through that Ψ . Since the algorithm maximizes $J_k = kJ(A, B, C) + J'$ in the unwrapped network for all k, it must maximize J(A, B, C). To see this, set the variables A, B, C, so that J_k is maximized. If J(A, B, C) is not maximized, then change A, B, and C to maximize J(A, B, C). This increases J_k by some quantity that is proportional to k. However, two of the variables that appear in copies of J(A, B, C) also appear in J'and thus J' might decrease in value. As long as J' decreases by some finite amount, we can increase J_k by increasing k sufficiently. As long as all Ψ 's are nonzero, J' which is proportional to $\log \Psi$, can change by at most some finite amount. Hence, for a network with a single loop, assuming that the message passing algorithm converges, it converges to the maximum a posteriori assignment.

9.10 Belief Update in Networks with a Single Loop

In the previous section, we showed that when the message passing algorithm converges, it correctly solves the MAP problem for graphs with a single loop. The message passing algorithm can also be used to obtain the correct answer for the marginalization problem. Consider a network consisting of a single loop with variables x_1, x_2, \ldots, x_n and evidence

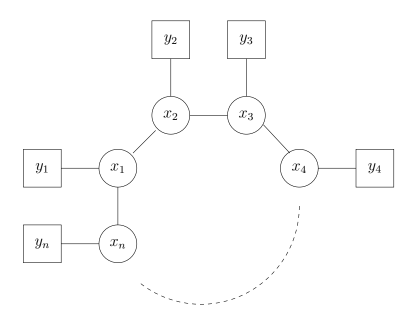


Figure 9.6: A Markov random field with a single loop.

 y_1, y_2, \ldots, y_n as shown in Figure 9.6. The x_i and y_i can be represented by vectors having a component for each value x_i can take on. To simplify the discussion assume the x_i take on values $1, 2, \ldots, m$.

Let m_i be the message sent from vertex i to vertex $i + 1 \mod n$. At vertex i + 1each component of the message m_i is multiplied by the evidence y_{i+1} and the constraint function Ψ . This is done by forming a diagonal matrix D_{i+1} where the diagonal elements are the evidence and then forming a matrix M_i whose jk^{th} element is $\Psi(x_{i+1} = j, x_i = k)$. The message m_{i+1} is $M_i D_{i+1} m_i$. Multiplication by the diagonal matrix D_{i+1} multiplies the components of the message m_i by the associated evidence. Multiplication by the matrix M_i multiplies each component of the vector by the appropriate value of Ψ and sums over the values producing the vector which is the message m_{i+1} . Once the message has travelled around the loop, the new message m'_1 is given by

$$m_1' = M_n D_1 M_{n-1} D_n \cdots M_2 D_3 M_1 D_2 m_1$$

Let $M = M_n D_1 M_{n-1} D_n \cdots M_2 D_3 M_1 D_2 m_1$. Assuming that *M*'s principle eigenvalue is unique, the message passing will converge to the principle vector of *M*. The rate of convergences depends on the ratio of the first and second eigenvalues.

An argument analogous to the above concerning the messages gong clockwise around the loop applies to messages moving counter clockwise around the loop. To obtain the estimate of the marginal probability $p(x_1)$, one multiples component wise the two messages arriving at x_1 along with the evidence y_1 . This estimate does not give the true marginal probability but the true marginal probability can be computed from the estimate and the rate of convergences by linear algebra.

9.11 Maximum Weight Matching

We have seen that the belief propagation algorithm converges to the correct solution in trees and graphs with a single cycle. It also correctly converges for a number of problems. Here we give one example, the maximum weight matching problem where there is a unique solution.

We apply the belief propagation algorithm to find the maximal weight matching (MWM) in a complete bipartite graph. If the MWM in the bipartite graph is unique, then the belief propagation algorithm will converge to it.

Let $G = (V_1, V_2, E)$ be a complete bipartite graph where $V_1 = \{a_1, \ldots, a_n\}, V_2 = \{b_1, \ldots, b_n\}$, and $(a_i, b_j) \in E$, $1 \leq i, j \leq n$. Let $\pi = \{\pi(1), \ldots, \pi(n)\}$ be a permutation of $\{1, \ldots, n\}$. The collection of edges $\{(a_1, b_{\pi(1)}), \ldots, (a_n, b_{\pi(n)})\}$ is called a *matching* which is denoted by π . Let w_{ij} be the weight associated with the edge (a_i, b_j) . The weight of the matching π is $w_{\pi} = \sum_{i=1}^{n} w_{i\pi(i)}$. The maximum weight matching π^* is $\pi^* = \arg \max_{\pi} w_{\pi}$

The first step is to create a factor graph corresponding to the MWM problem. Each edge of the bipartite graph is represented by a variable c_{ij} which takes on the value zero or one. The value one means that the edge is present in the matching, the value zero means that the edge is not present in the matching. A set of constraints is used to force the set of edges to be a matching. The constraints are of the form $\sum_{j} c_{ij} = 1$ and $\sum_{i} c_{ij} = 1$. Any 0,1 assignment to the variables c_{ij} that satisfies all of the constraints defines a matching. In addition, we have constraints for the weights of the edges.

We now construct a factor graph, a portion of which is shown in Fig. 9.8. Associated with the factor graph is a function $f(c_{11}, c_{12}, ...)$ consisting of a set of terms for each c_{ij} enforcing the constraints and summing the weights of the edges of the matching. The terms for c_{12} are

$$-\lambda \left| \left(\sum_{i} c_{i2} \right) - 1 \right| - \lambda \left| \left(\sum_{j} c_{1j} \right) - 1 \right| + w_{12}c_{12}$$

where λ is a large positive number used to enforce the constraints when we maximize the function. Finding the values of c_{11}, c_{12}, \ldots that maximize f finds the maximum weighted matching for the bipartite graph.

If the factor graph was a tree, then the message from a variable node x to its parent is a message g(x) that gives the maximum value for the sub tree for each value of x. To compute g(x), one sums all messages into the node x. For a constraint node, one sums all messages from sub trees and maximizes the sum over all variables except the variable of the parent node subject to the constraint. The message from a variable x consists of two pieces of information, the value p(x = 0) and the value p(x = 1). This information can be encoded into a linear function of x.

$$[p(x = 1) - p(x = 0)]x + p(x = 0)$$

Thus, the messages are of the form ax + b. To determine the MAP value of x once the algorithm converges, sum all messages into x and take the maximum over x=1 and x=0 to determine the value for x. Since the arg maximum of a linear form ax+b depends only on whether a is positive or negative and since maximizing the output of a constraint depends only on the coefficient of the variable, we can send messages consisting of just the variable coefficient.

To calculate the message to c_{12} from the constraint that node b_2 has exactly one neighbor, add all the messages that flow into the constraint node from the c_{i2} , $i \neq 1$ nodes and maximize subject to the constraint that exactly one variable has value one. If $c_{12} = 0$, then one of c_{i2} , $i \neq 1$, will have value one and the message is $\max_{i\neq 1} \alpha(i, 2)$. If $c_{12} = 1$, then the message is zero. Thus, we get

$$-\max_{i\neq 1}\alpha\left(i,2\right)x + \max_{i\neq 1}\alpha\left(i,2\right)$$

and send the coefficient $-\max_{i\neq 1} \alpha(i,2)$. This means that the message from c_{12} to the other constraint node is $\beta(1,2) = w_{12} - \max_{i\neq 1} \alpha(i,2)$.

The alpha message is calculated in a similar fashion. If $c_{12} = 0$, then one of c_{1j} will have value one and the message is $\max_{j \neq 1} \beta(1, j)$. If $c_{12} = 1$, then the message is zero. Thus, the coefficient $-\max_{j \neq 1} \alpha(1, j)$ is sent. This means that $\alpha(1, 2) = w_{12} - \max_{j \neq 1} \alpha(1, j)$.

To prove convergence, we enroll the constraint graph to form a tree with a constraint node as the root. In the enrolled graph a variable node such as c_{12} will appear a number of times which depends on how deep a tree is built. Each occurrence of a variable such as c_{12} is deemed to be a distinct variable.

Lemma 9.3 If the tree obtained by unrolling the graph is of depth k, then the messages to the root are the same as the messages in the constraint graph after k-iterations.

Proof: Straight forward.

Define a matching in the tree to be a set of vertices so that there is exactly one variable node of the match adjacent to each constraint. Let Λ denote the vertices of the matching. Heavy circles represent the nodes of the above tree that are in the matching Λ .

Let Π be the vertices corresponding to maximum weight matching edges in the bipartite graph. Recall that vertices in the above tree correspond to edges in the bipartite

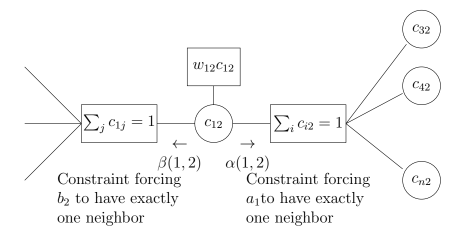


Figure 9.7: Portion of factor graph for the maximum weight matching problem.

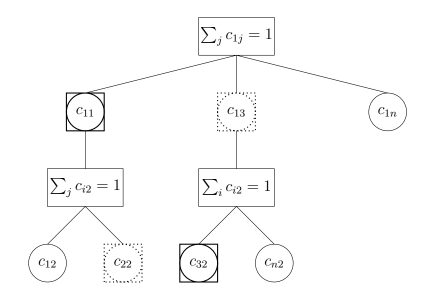


Figure 9.8: Tree for MWM problem.

graph. The vertices of Π are denoted by dotted circles in the above tree.

Consider a set of trees where each tree has a root that corresponds to one of the constraints. If the constraint at each root is satisfied by the edge of the MWM, then we have found the MWM. Suppose that the matching at the root in one of the trees disagrees with the MWM. Then there is an alternating path of vertices of length 2k consisting of vertices corresponding to edges in Π and edges in Λ . Map this path onto the bipartite graph. In the bipartite graph the path will consist of a number of cycles plus a simple path. If k is large enough there will be a large number of cycles since no cycle can be of length more than 2n. Let m be the number of cycles. Then $m \geq \frac{2k}{2n} = \frac{k}{n}$.

Let π^* be the MWM in the bipartite graph. Take one of the cycles and use it as an alternating path to convert the MWM to another matching. Assuming that the MWM is unique and that the next closest matching is ε less, $W_{\pi^*} - W_{\pi} > \varepsilon$ where π is the new matching.

Consider the tree matching. Modify the tree matching by using the alternating path of all cycles and the left over simple path. The simple path is converted to a cycle by adding two edges. The cost of the two edges is at most $2w^*$ where w^* is the weight of the maximum weight edge. Each time we modify Λ by an alternating cycle, we increase the cost of the matching by at least ε . When we modify Λ by the left over simple path, we increase the cost of the tree matching by $\varepsilon - 2w^*$ since the two edges that were used to create a cycle in the bipartite graph are not used. Thus

weight of Λ - weight of $\Lambda' \geq \frac{k}{n}\varepsilon - 2w*$

which must be negative since Λ' is optimal for the tree. However, if k is large enough this becomes positive, an impossibility since Λ' is the best possible. Since we have a tree, there can be no cycles, as messages are passed up the tree, each sub tree is optimal and hence the total tree is optimal. Thus the message passing algorithm must find the maximum weight matching in the weighted complete bipartite graph assuming that the maximum weight matching is unique. Note that applying one of the cycles that makes up the alternating path decreased the bipartite graph match but increases the value of the tree. However, it does not give a higher tree matching, which is not possible since we already have the maximum tree matching. The reason for this is that the application of a single cycle does not result in a valid tree matching. One must apply the entire alternating path to go from one matching to another.

9.12 Warning Propagation

Significant progress has been made using methods similar to belief propagation in finding satisfying assignments for 3-CNF formulas. Thus, we include a section on a version of belief propagation, called warning propagation, that is quite effective in finding assignments. Consider a factor graph for a SAT problem. Index the variables by i, j, and

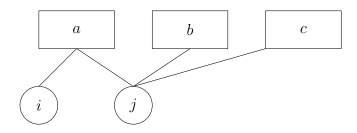


Figure 9.9: warning propagation

k and the factors by a, b, and c. Factor a sends a message m_{ai} to each variable i that appears in the factor a called a warning. The warning is 0 or 1 depending on whether or not factor a believes that the value assigned to i is required for a to be satisfied. A factor a determines the warning to send to variable i by examining all warnings received by other variables in factor a from factors containing them.

For each variable j, sum the warnings from factors containing j that warn j to take value T and subtract the warnings that warn j to take value F. If the difference says that j should take value T or F and this value for variable j does not satisfy a, and this is true for all j, then a sends a warning to i that the value of variable i is critical for factor a.

Start the warning propagation algorithm by assigning 1 to a warning with probability 1/2. Iteratively update the warnings. If the warning propagation algorithm converges, then compute for each variable *i* the local field h_i and the contradiction number c_i . The local field h_i is the number of clauses containing the variable *i* that sent messages that *i* should take value T minus the number that sent messages that *i* should take value F. The contradiction number c_i is 1 if variable *i* gets conflicting warnings and 0 otherwise. If the factor graph is a tree, the warning propagation algorithm converges. If one of the warning messages is one, the problem is unsatisfiable; otherwise it is satisfiable.

9.13 Correlation Between Variables

In many situations one is interested in how the correlation between variables drops off with some measure of distance. Consider a factor graph for a 3-CNF formula. Measure the distance between two variables by the shortest path in the factor graph. One might ask if one variable is assigned the value true, what is the percentage of satisfying assignments of the 3-CNF formula in which the second variable also is true. If the percentage is the same as when the first variable is assigned false, then we say that the two variables are uncorrelated. How difficult it is to solve a problem is likely to be related to how fast the correlation decreases with distance.

Another illustration of this concept is in counting the number of perfect matchings in a graph. One might ask what is the percentage of matching in which some edge is present and ask how correlated this percentage is with the presences or absence of edges at some distance d. One is interested in whether the correlation drops off with distance. To explore this concept we consider the Ising model studied in physics.

The Ising or ferromagnetic model is a pairwise random Markov field. The underlying graph, usually a lattice, assigns a value of ± 1 , called spin, to the variable at each vertex. The probability (Gibbs measure) of a given configuration of spins is proportional to $exp(\beta \sum_{(i,j)\in E} x_i x_j) = \prod_{(i,j)\in E} e^{\beta x_i x_j}$ where $x_i = \pm 1$ is the value associated with vertex *i*.

Thus

$$p(x_1, x_2, \dots, x_n) = \frac{1}{Z} \prod_{(i,j) \in E} exp(\beta x_i x_j) = \frac{1}{Z} e^{\beta \sum_{(i,j) \in E} x_i x_j}$$

0

where Z is a normalization constant.

The value of the summation is simply the difference in the number of edges whose vertices have the same spin minus the number of edges whose vertices have opposite spin. The constant β is viewed as inverse temperature. High temperature corresponds to a low value of β and low temperature corresponds to a high value of β . At high temperature, low β , the spins of adjacent vertices are uncorrelated whereas at low temperature adjacent vertices have identical spins. The reason for this is that the probability of a configuration is proportional to $e^{\beta \sum_{i < j} x_i x_j}$. As β is increased, for configurations with a large number of edges whose vertices have identical spins, $e^{\beta \sum_{i < j} x_i x_j}$ increases more than for configurations whose edges have vertices with non identical spins. When the normalization constant $\frac{1}{Z}$ is adjusted for the new value of β , the highest probability configurations are those where adjacent vertices have identical spins.

Given the above probability distribution, what is the correlation between two variables x_i and x_j . To answer this question, consider the probability that x_i equals plus one as a function of the probability that x_j equals plus one. If the probability that x_i equals plus one is $\frac{1}{2}$ independent of the value of the probability that x_j equals plus one, we say the values are uncorrelated.

Consider the special case where the graph G is a tree. In this case a phase transition occurs at $\beta_0 = \frac{1}{2} \ln \frac{d+1}{d-1}$ where d is the degree of the tree. For a sufficiently tall tree and for $\beta > \beta_0$, the probability that the root has value +1 is bounded away from $\frac{1}{2}$ and depends on whether the majority of leaves have value +1 or -1. For $\beta < \beta_0$ the probability that the root has value +1 is $\frac{1}{2}$ independent of the values at the leaves of the tree.

Consider a height one tree of degree d. If i of the leaves have spin +1 and d - i have spin -1, then the probability of the root having spin +1 is proportional to

$$e^{i\beta - (d-i)\beta} = e^{(2i-d)\beta}.$$

If the probability of a leaf being +1 is p, then the probability of i leaves being +1 and d-i being -1 is

$$\binom{d}{i} p^i \left(1 - p\right)^{d-i}$$

Thus, the probability of the root being +1 is proportional to

$$A = \sum_{i=1}^{d} {\binom{d}{i}} p^{i} (1-p)^{d-i} e^{(2i-d)\beta} = e^{-d\beta} \sum_{i=1}^{d} {\binom{d}{i}} \left(p e^{2\beta} \right)^{i} (1-p)^{d-i} = e^{-d\beta} \left[p e^{2\beta} + 1 - p \right]^{d-i}$$

and the probability of the root being -1 is proportional to

$$B = \sum_{i=1}^{d} {\binom{d}{i}} p^{i} (1-p)^{d-i} e^{-(2i-d)\beta}$$

= $e^{-d\beta} \sum_{i=1}^{d} {\binom{d}{i}} p^{i} \left[(1-p) e^{-2(i-d)\beta} \right]$
= $e^{-d\beta} \sum_{i=1}^{d} {\binom{d}{i}} p^{i} \left[(1-p) e^{2\beta} \right]^{d-i}$
= $e^{-d\beta} \left[p + (1-p) e^{2\beta} \right]^{d}$.

The probability of the root being +1 is

$$q = \frac{A}{A+B} = \frac{\left[pe^{2\beta} + 1 - p\right]^d}{\left[pe^{2\beta} + 1 - p\right]^d + \left[p + (1-p)e^{2\beta}\right]^d} = \frac{C}{D}$$

where

$$C = \left[p e^{2\beta} + 1 - p \right]^d$$

and

$$D = [pe^{2\beta} + 1 - p]^{d} + [p + (1 - p)e^{2\beta}]^{d}.$$

At high temperature, low β , the probability q of the root of the height one tree being +1 in the limit as β goes to zero is

$$q = \frac{p+1-p}{[p+1-p] + [p+1-p]} = \frac{1}{2}$$

independent of p. At low temperature, high β ,

$$q \approx \frac{p^d e^{2\beta d}}{p^d e^{2\beta d} + (1-p)^d e^{2\beta d}} = \frac{p^d}{p^d + (1-p)^d} = \begin{cases} 0 & p = 0\\ 1 & p = 1 \end{cases}.$$

q goes from a low probability of +1 for p below 1/2 to high probability of +1 for p above 1/2.

Now consider a very tall tree. If the p is the probability that a root has value +1, we can iterate the formula for the height one tree and observe that at low temperature the probability of the root being one converges to some value. At high temperature, the probability of the root being one is 1/2 independent of p. See Figure 9.10. At the phase transition, the slope of q at p=1/2 is one.

Now the slope of the probability of the root being 1 with respect to the probability of a leaf being 1 in this height one tree is

$$\frac{\partial q}{\partial p} = \frac{D\frac{\partial C}{\partial p} - C\frac{\partial D}{\partial p}}{D^2}$$

Since the slope of the function q(p) at p=1/2 when the phase transition occurs is one, we can solve $\frac{\partial q}{\partial p} = 1$ for the value of β where the phase transition occurs. First, we show that $\left. \frac{\partial D}{\partial p} \right|_{p=\frac{1}{2}} = 0.$

$$D = \left[p e^{2\beta} + 1 - p \right]^d + \left[p + (1 - p) e^{2\beta} \right]^d$$
$$\frac{\partial D}{\partial p} = d \left[p e^{2\beta} + 1 - p \right]^{d-1} \left(e^{2\beta} - 1 \right) + d \left[p + (1 - p) e^{2\beta} \right]^{d-1} \left(1 - e^{2\beta} \right)$$
$$\frac{\partial D}{\partial p} \Big|_{p=\frac{1}{2}} = \frac{d}{2^{d-1}} \left[e^{2\beta} + 1 \right]^{d-1} \left(e^{2\beta} - 1 \right) + \frac{d}{2^{d-1}} \left[1 + e^{2\beta} \right]^{d-1} \left(1 - e^{2\beta} \right) = 0$$

Then

$$\begin{aligned} \frac{\partial q}{\partial p}\Big|_{p=\frac{1}{2}} &= \frac{D\frac{\partial C}{\partial p} - C\frac{\partial D}{\partial p}}{D^2} \Big|_{p=\frac{1}{2}} = \frac{\frac{\partial C}{\partial p}}{D} \Big|_{p=\frac{1}{2}} = \frac{d\left[pe^{2\beta} + 1 - p\right]^{d-1} \left(e^{2\beta} - 1\right)}{\left[pe^{2\beta} + 1 - p\right]^d + \left[p + (1 - p)e^{2\beta}\right]^d} \Big|_{p=\frac{1}{2}} \\ &= \frac{d\left[\frac{1}{2}e^{2\beta} + \frac{1}{2}\right]^{d-1} \left(e^{2\beta} - 1\right)}{\left[\frac{1}{2}e^{2\beta} + \frac{1}{2}\right]^d + \left[\frac{1}{2} + \frac{1}{2}e^{2\beta}\right]^d} = \frac{d\left(e^{2\beta} - 1\right)}{1 + e^{2\beta}}\end{aligned}$$

Setting

$$\frac{d(e^{2\beta} - 1)}{1 + e^{2\beta}} = 1$$

And solving for β yields

$$d(e^{2\beta} - 1) = 1 + e^{2\beta}$$
$$e^{2\beta} = \frac{d+1}{d-1}$$
$$\beta = \frac{1}{2} \ln \frac{d+1}{d-1}$$

To complete the argument, we need to show that q is a monotonic function of p. To see this, write $q = \frac{1}{1 + \frac{B}{A}}$. A is a monotonically increasing function of p and B is monotonically

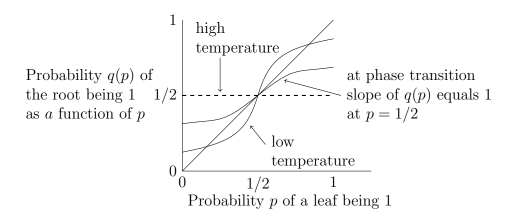


Figure 9.10: Shape of q as a function of p for the height one tree and three values of β corresponding to low temperature, the phase transition temperature, and high temperature.

decreasing. From this it follows that q is monotonically increasing.

In the iteration going from p to q, we do not get the true marginal probabilities at each level since we ignored the effect of the portion of the tree above. However, when we get to the root, we do get the true marginal for the root. To get the true marginal's for the interior nodes we need to send messages down from the root.

Note: The joint probability distribution for the tree is of the form $e^{\beta \sum_{(ij) \in E} x_i x_j} = \prod_{(i,j) \in E} e^{\beta x_i x_j}$. Suppose x_1 has value 1 with probability p. Then define a function φ , called evidence, such that

$$\varphi(x_1) = \begin{cases} p & \text{for } \mathbf{x}_1 = 1\\ 1 - p & \text{for } \mathbf{x}_1 = -1\\ = \left(p - \frac{1}{2}\right)x_1 + \frac{1}{2} \end{cases}$$

and multiply the joint probability function by φ . Note, however, that the marginal probability of x_1 is not p. In fact, it may be further from p after multiplying the conditional probability function by the function φ .

9.14 Exercises

Exercise 9.1 Find a nonnegative factorization of the matrix

$$A = \begin{pmatrix} 4 & 6 & 5\\ 1 & 2 & 3\\ 7 & 10 & 7\\ 6 & 8 & 4\\ 6 & 10 & 11 \end{pmatrix}$$

Indicate the steps in your method and show the intermediate results.

Exercise 9.2 Find a nonnegative factorization of each of the following matrices.

(1)	$ \begin{pmatrix} 10 \\ 2 \\ 8 \\ 7 \\ 5 \\ 1 \\ 2 \end{pmatrix} $	$\frac{1}{7}$	3 13 11 11	$14 \\ 3 \\ 11 \\ 10 \\ 6 \\ 1$	13 1 11 7 11 3 2				(2)	2	2 1 1 3	4	$ \begin{array}{r} 14 \\ 4 \\ 2 \\ 8 \\ 16 \\ 6 \end{array} $	$ \begin{array}{c} 17 \\ 6 \\ 4 \\ 3 \\ 10 \\ 18 \\ 7 \end{array} \right) $		
(3)	$\begin{pmatrix} 4 \\ 13 \\ 15 \\ 7 \\ 1 \\ 5 \\ 3 \end{pmatrix}$	$\begin{array}{c} 4 \\ 16 \\ 24 \\ 16 \\ 4 \\ 8 \\ 12 \end{array}$	$3 \\ 13 \\ 21 \\ 15 \\ 4 \\ 7 \\ 12$	$ \begin{array}{r} 10 \\ 12 \\ 6 \\ 1 \end{array} $	$ \begin{array}{c} 1 \\ 5 \\ 9 \\ 7 \\ 2 \\ 3 \\ 6 \end{array} $	$3 \\ 13 \\ 21 \\ 15 \\ 4 \\ 7 \\ 12$	$\begin{array}{c} 4 \\ 14 \\ 18 \\ 10 \\ 2 \\ 6 \\ 6 \\ \end{array}$	$ \begin{array}{c} 3 \\ 10 \\ 12 \\ 6 \\ 1 \\ 4 \\ 3 \end{array} \right) $	(4)	$\begin{pmatrix} 1\\9\\6\\3 \end{pmatrix}$	$egin{array}{c} 1 \\ 9 \\ 6 \\ 3 \end{array}$	$3 \\ 9 \\ 12 \\ 3$	4 12 16 4	4 9 15 3	4 9 15 3	$\begin{pmatrix} 1\\3\\4\\1 \end{pmatrix}$

Exercise 9.3 Consider the matrix A that is the product of nonnegative matrices B and C. (12, 22, 11, 25) (12, 1)

/12					(10)					
19	20	13	48		1	9	$\begin{pmatrix} 1 \\ 2 \end{pmatrix}$	2	4	3
11	14	16	29	=	3	4	$\backslash 2$	2	1	5/
14	16	14	36/		2	6/	,			,

Which rows of A are approximate positive linear combinations of other rows of A? Find an approximate nonnegative factorization of A

Exercise 9.4 What is the probability of heads occurring after a sufficiently long sequence of transitions in Viterbi algorithm example of the most likely sequence of states?

Exercise 9.5 Find optimum parameters for a three state HMM and given output sequence. Note the HMM must have a strong signature in the output sequence or we probably will not be able to find it. The following example may not be good for that reason.

	1	2	3		A	В
1	$\frac{1}{2}$	$\frac{1}{4}$	$\frac{1}{4}$	1	$\frac{3}{4}$	$\frac{1}{4}$
2	$\frac{1}{4}$	$\frac{1}{4}$	$\frac{1}{2}$	2	$\frac{1}{4}$	$\frac{3}{4}$
3	$\frac{1}{3}$	$\frac{1}{3}$	$\frac{1}{3}$	3	$\frac{1}{3}$	$\frac{2}{3}$

Exercise 9.6 In the Ising model for a tree of degree one, a chain of vertices, is there a phase transition where the correlation between the value at the root and the value at the leaves becomes independent? Work out mathematical what happens.

Exercise 9.7 For a Boolean function in CNF the marginal probability gives the number of satisfiable assignments with x_1 .

How does one obtain the number of satisfying assignments for a 2-CNF formula? Not completely related to first sentence.

10 Other Topics

10.1 Rankings

Ranking is important. We rank movies, restaurants, students, web pages, and many other items. Ranking has become a multi-billion dollar industry as organizations try to raise the position of their web pages in the display of web pages returned by search engines to relevant queries. Developing a method of ranking that is not manipulative is an important task.

A ranking is a complete ordering in the sense that for every pair of items a and b, either a is preferred to b or b is preferred to a. Furthermore, a ranking is transitive in that a > b and b > c implies a > c.

One problem of interest in ranking is that of combining many individual rankings into one global ranking. However, merging ranked lists is nontrivial as the following example illustrates.

Example: Suppose there are three individuals who rank items a, b, and c as illustrated in the following table.

individual	first item	second item	third item
1	a	b	С
2	b	С	a
3	С	a	b

Suppose our algorithm tried to rank the items by first comparing a to b and then comparing b to c. In comparing a to b, two of the three individuals prefer a to b and thus we conclude a is preferable to b. In comparing b to c, again two of the three individuals prefer b to c and we conclude that b is preferable to c. Now by transitivity one would expect that the individuals would prefer a to c, but such is not the case, only one of the individuals prefers a to c and thus c is preferable to a. We come to the illogical conclusion that a is preferable to b, b is preferable to c, and c is preferable to a.

Suppose there are a number of individuals or voters and a set of candidates to be ranked. Each voter produces a ranked list of the candidates. From the set of ranked lists can one construct a single ranking of the candidates? Assume the method of producing a global ranking is required to satisfy the following three axioms.

Nondictatorship – The algorithm cannot always simply select one individual's ranking.

Unanimity – If every individual prefers a to b, then the global ranking must prefer a to b.

Independent of irrelevant alternatives – If individuals modify their rankings but keep the order of a and b unchanged, then the global order of a and b should not change.

Arrow showed that no ranking algorithm exists satisfying the above axioms.

Theorem 10.1 (*Arrow*) Any algorithm for creating a global ranking from individual rankings of three or more elements in which the global ranking satisfies unanimity and independence of irrelevant alternatives is a dictatorship.

Proof: Let a, b, and c be distinct items. Consider a set of rankings in which each individual ranks b either first or last. Some individuals may rank b first and others may rank b last. For this set of rankings, the global ranking must put b first or last. Suppose to the contrary that b is not first or last in the global ranking. Then there exist a and c where the global ranking puts a > b and b > c. By transitivity, the global ranking puts a > c. Note that all individuals can move c above a without affecting the order of b and a or the order of b and c since b was first or last on each list. Thus, by independence of irrelevant alternatives, the global ranking would continue to rank a > b and b > c even if all individuals moved c above a since that would not change the individuals relative order of a and b or the individuals relative order of b and c. But then by unanimity, the global ranking would need to put c > a, a contradiction. We conclude that the global ranking puts b first or last.

Consider a set of rankings in which every individual ranks b last. By unanimity, the global ranking must also rank b last. Let the individuals, one by one, move b from bottom to top leaving the other rankings in place. By unanimity, the global ranking must eventually move b from the bottom all the way to the top. When b first moves, it must move all the way to the top by the previous argument. Let v be the first individual whose change causes the global ranking of b to change.

We now argue that v is a dictator. First, we argue that v is a dictator for any pair ac not involving b. We will refer to three rankings of v (see Figure 10.1). The first ranking of v is the ranking prior to v moving b from the bottom to the top and the second is the ranking just after v has moved b to the top. Choose any pair ac where a is above c in v's ranking. The third ranking of v is obtained by moving a above b in the second ranking so that a > b > c in v's ranking. By independence of irrelevant alternatives, the global ranking after v has switched to the third ranking puts a > b since all individual ab votes are the same as just before v moved b to the top of his ranking. At that time the global ranking placed a > b. Similarly b > c in the global ranking since all individual bc votes are the same as just after v moved b to the top causing b to move to the top in the global ranking. By transitivity the global ranking must put a > c and thus the global ranking of a and c agrees with v.

Now all individuals except v can modify their rankings arbitrarily while leaving b in its extreme position and by independence of irrelevant alternatives, this does not affect the

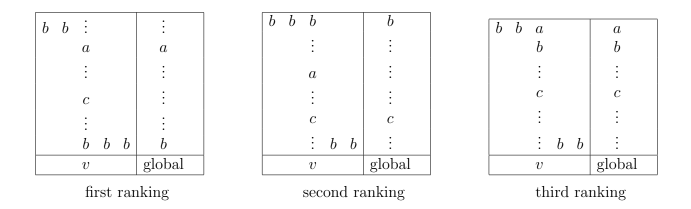


Figure 10.1: The three rankings that are used in the proof of Theorem 10.1.

global ranking of a > b or of b > c. Thus, by transitivity this does not affect the global ranking of a and c. Next, all individuals except v can move b to any position without affecting the global ranking of a and c.

At this point we have argued that independent of other individuals' rankings, the global ranking of a and c will agree with v's ranking. Now v can change its ranking arbitrarily, provided it maintains the order of a and c, and by independence of irrelevant alternatives the global ranking of a and c will not change and hence will agree with v. Thus, we conclude that for all a and c, the global ranking agrees with v independent of the other rankings except for the placement of b. But other rankings can move b without changing the global order of other elements. Thus, v is a dictator for the ranking of any pair of elements not involving b.

Note that v changed the relative order of a and b in the global ranking when it moved b from the bottom to the top in the previous argument. We will use this in a moment.

The individual v is also a dictator over every pair ab. Repeat the construction showing that v is a dictator for every pair ac not involving b only this time place c at the bottom. There must be an individual v_c who is a dictator for any pair such as ab not involving c. Since both v and v_c can affect the global ranking of a and b independent of each other, it must be that v_c is actually v. Thus, the global ranking agrees with v no matter how the other voters modify their rankings.

10.2 Hare System for Voting

One voting system would be to have everyone vote for their favorite candidate. If some candidate receives a majority of votes, he or she is declared the winner. If no candidate receives a majority of votes, the candidate with the fewest votes is dropped from the slate and the process is repeated. The Hare system implements this method by asking each voter to rank all the candidates. Then one counts how many voters ranked each candidate as number one. If no candidate receives a majority, the candidate with the fewest number one votes is dropped from each voters ranking. If the dropped candidate was number one on some voters list, then the number two candidate becomes that voter's number one choice. The process of counting the number one rankings is then repeated.

Although the Hare system is widely used it fails to satisfy Arrow' axioms as all voting systems must. Consider the following situation in which there are 21 voters that fall into four categories. Voters within a category rank individuals in the same order.

Category	Number of voters in category	Preference order
1	7	abcd
2	6	bacd
3	5	cbad
4	3	dcba

The Hare system would first eliminate d since d gets only three rank one votes. Then it would eliminate b since b gets only six rank one votes whereas a gets seven and c gets eight. At this point a is declared the winner since a has thirteen votes to c's eight votes.

Now assume that Category 4 voters who prefer b to a move a up to first place. Then the election proceeds as follows. In round one, d is eliminated since it gets no rank one votes. Then c with five votes is eliminated and b is declared the winner with 11 votes. Note that by moving a up, category 4 voters were able to deny a the election and get bto win, whom they prefer over a.

10.3 Compressed Sensing and Sparse Vectors

Given a function x(t), one can represent the function by the composition of sinusoidal functions. Basically one is representing the time function by its frequency components. The transformation from the time representation of a function to it frequency representation is accomplished by a Fourier transform. The Fourier transform of a function x(t)is given by

$$f(\omega) = \int x(t) e^{-2\pi\omega t} dt$$

Converting the frequency representation back to the time representation is done by the inverse Fourier transformation

$$x(t) = \int f(\omega)e^{-2\pi\omega t}d\omega$$

In the discrete case, $\mathbf{x} = [x_0, x_1, \dots, x_{n-1}]$ and $\mathbf{f} = [f_0, f_1, \dots, f_{n-1}]$. The Fourier transform and its inverse are $\mathbf{f} = A\mathbf{x}$ with $a_{ij} = \omega^{ij}$ where ω is the principle n^{th} root of unity.

There are many other transforms such as the Laplace, wavelets, chirplets, etc. In fact, any nonsingular $n \times n$ matrix can be used as a transform.

If one has a discrete time sequence \mathbf{x} of length n, the Nyquist theorem states that n coefficients in the frequency domain are needed to represent the signal \mathbf{x} . However, if the signal \mathbf{x} has only s nonzero elements, even though one does not know which elements they are, one can recover the signal by randomly selecting a small subset of the coefficients in the frequency domain. It turns out that one can reconstruct sparse signals with far fewer samples than one might suspect and an area called compressed sampling has emerged with important applications.

Motivation

Let A be an $n \times d$ matrix with n much smaller than d whose elements are generated by independent, zero mean, unit variance, Gaussian processes. Let **x** be a sparse d-dimensional vector with at most s nonzero coordinates, $s \ll d$. **x** is called the signal and A is the "measurement" matrix. What we measure are the components of the n dimensional vector A**x**. We ask if we can recover the signal **x** from measurements A**x**, where the number n of measurements is much smaller than the dimension d? We have two advantages over an arbitrary system of linear equations. First, the solution **x** is known to be sparse and second we have the choice of the measurement matrix A.

In many applications, the signal is sparse in either the time domain or the frequency domain. For images, it is often the case that in the frequency domain very few frequencies have significant amplitude. If we zero out small frequency amplitudes, we get a sparse frequency representation of the signal. It is wasteful to measure each of the *d* components of the signal \mathbf{x} , most of which are zero. Instead, we measure *n* linear combinations of components, the linear combinations form *A*. In applications, we choose the matrix *A*. A usual choice is a matrix whose entries are independent zero mean, unit variance Gaussian random variables. Since we have no control over the signal, our system needs to recover any signal. We will show that *n* needs to depend essentially only on *s*, not on *d*.

10.3.1 Unique Reconstruction of a Sparse Vector

A vector is said to be s-sparse if it has at most s nonzero elements. Let **x** be a ddimensional, s-sparse vector with $s \ll d$. Consider solving $A\mathbf{x} = \mathbf{b}$ for **x** where A is an $n \times d$ matrix with $n \ll d$. The set of solutions to $A\mathbf{x} = \mathbf{b}$ is a subspace. However, if we restrict ourselves to sparse solutions, under certain conditions on A there is a unique ssparse solution. Suppose that there were two s-sparse solutions, \mathbf{x}_1 and \mathbf{x}_2 . Then $\mathbf{x}_1 - \mathbf{x}_2$

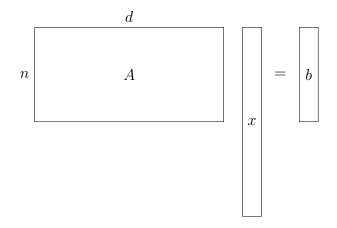


Figure 10.2: $A\mathbf{x} = \mathbf{b}$ has a vector space of solutions but possibly only one sparse solution.

would be a 2s-sparse solution to the homogeneous system $A\mathbf{x} = \mathbf{0}$. A 2s-sparse solution to the homogeneous equation $A\mathbf{x} = \mathbf{0}$ requires that some 2s columns of A be linearly dependent. Unless A has 2s linearly dependent columns there can be only one s-sparse solution.

Now suppose n is order s^2 and we pick an $n \times d$ matrix A with random independent zero mean, unit variance Gaussian entries. Take any subset of 2s columns of A. Since we have already seen in Chapter 2 that each of these 2s vectors is likely to be essentially orthogonal to the space spanned by the previous vectors, the sub-matrix is unlikely to be singular. This intuition can be made rigorous.

To find a sparse solution to $A\mathbf{x} = \mathbf{b}$, one would like to minimize the zero norm $\|\mathbf{x}\|_0$ over $\{\mathbf{x}|A\mathbf{x} = \mathbf{b}\}$. This is a computationally hard problem. There are techniques to minimize a convex function over a convex set. But $\||\mathbf{x}\||_0$ is not a convex function. With no further hypotheses, it is NP-hard. With this in mind, we use the one norm as a proxy for the zero norm and minimize the one norm $\|\mathbf{x}\|_1$ over $\{\mathbf{x}|A\mathbf{x} = \mathbf{b}\}$. Although this problem appears to be nonlinear, it can be solved by linear programming by writing $\mathbf{x} = \mathbf{u} - \mathbf{v}$, $\mathbf{u} \ge 0$, and $\mathbf{v} \ge 0$, and then minimizing the linear function $\sum_i u_i + \sum_i v_i$ subject to $A\mathbf{u}$ - $A\mathbf{v}=\mathbf{b}$, $\mathbf{u} \ge 0$, and $\mathbf{v} \ge 0$.

Under what conditions will minimizing $\|\mathbf{x}\|_1$ over $\{\mathbf{x}|A\mathbf{x} = \mathbf{b}\}$ recover the *s*-sparse solution to $A\mathbf{x}=\mathbf{b}$? If $g(\mathbf{x})$ is a convex function, then any local minimum of g is a global minimum. If $g(\mathbf{x})$ is differentiable at its minimum, the gradient ∇g must be zero there. However, the 1-norm is not differentiable at its minimum. Thus, we introduce the concept of a subgradient of a convex function. Where the function is differentiable the subgradient is the gradient. Where the function is not differentiable, the sub gradient is any line touching the function at the point that lies totally below the function. See Figure 10.3.

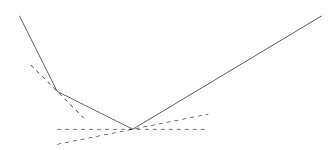


Figure 10.3: Some subgradients for a function that is not everywhere differentiable.

Subgradients are defined as follows. A subgradient of a function g at a point \mathbf{x}_0 , is a vector $\nabla g(\mathbf{x}_0)$ satisfying $g(\mathbf{x}_0 + \Delta \mathbf{x}) \ge g(\mathbf{x}_0) + (\nabla g)^T \Delta \mathbf{x}$ for any vector $\Delta \mathbf{x}$. A point is a minimum for a convex function if there is a subgradient at that point with slope zero.

Consider the function $||x||_1$, where x is a real variable. For x < 0, the subgradient equals the gradient and has value -1. For x > 0, the subgradient equals the gradient and has value 1. At x = 0, the subgradient can be any value in the range [-1,1]. The following proposition generalizes this example to the 1-norm function in d-space.

Proposition 10.2 A vector \mathbf{v} is a subgradient of the 1-norm function $||\mathbf{x}||_1$ at \mathbf{x} if and only if it satisfies the three conditions below:

- 1. $v_i = -1$ for all *i* in I_1 where, $I_1 = \{i | x_i < 0\}$,
- 2. $v_i = 1$ for all *i* in I_2 where, $I_2 = \{i | x_i > 0\}$, and
- 3. v_i in [-1, 1] for all *i* in I_3 where, $I_3 = \{i | x_i = 0\}$.

Proof: It is easy to see that for any vector **y**,

$$||\mathbf{x} + \mathbf{y}||_1 - ||\mathbf{x}||_1 \ge -\sum_{i \in I_1} y_i + \sum_{i \in I_2} y_i + \sum_{i \in I_3} |y_i|.$$

If *i* is in I_1 , x_i is negative. If y_i is also negative, then $||x_i + y_i||_1 = ||x_i||_1 + ||y_i||_1$ and thus $||x_i + y_i||_1 - ||x_i||_1 = ||y_i||_1 = -y_i$. If y_i is positive and less than $||x_i||_1$, then $||x_i + y_i||_1 = ||x_i|| - ||x_i||_1 = -y_i$. If y_i is positive and greater than or equal to $||x_i||_1$, then $||x_i + y_i||_1 = y_i - ||x_i||_1$ and thus $||x_i + y_i||_1 - ||x_i||_1 = y_i - 2||x_i||_1 \ge -y_i$. Similar reasoning establishes the case for *i* in I_2 or I_3 .

If \mathbf{v} satisfies the conditions in the proposition, then $||\mathbf{x}+\mathbf{y}||_1 \ge ||\mathbf{x}||_1 + \mathbf{v}^T \mathbf{y}$ as required. Now for the converse, suppose that \mathbf{v} is a subgradient. Consider a vector \mathbf{y} that is zero in all components except the first and y_1 is nonzero with $y_1 = \pm \varepsilon$ for a small $\varepsilon > 0$. If $1 \in I_1$, then $||\mathbf{x}+\mathbf{y}||_1 - ||\mathbf{x}||_1 = -y_1$ which implies that $-y_1 \ge v_1 y_1$. Choosing $y_1 = \varepsilon$,

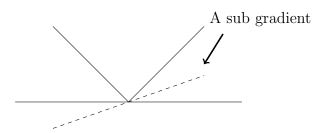


Figure 10.4: Illustration of a subgradient for $|\mathbf{x}|_1$ at $\mathbf{x} = 0$

gives $-1 \ge v_1$ and choosing $y_1 = -\varepsilon$, gives $-1 \le v_1$. So $v_1 = -1$. Similar reasoning gives the second condition. For the third condition, choose *i* in I_3 and set $y_i = \pm \varepsilon$ and argue similarly.

To characterize the value of \mathbf{x} that minimizes $\|\mathbf{x}\|_1$ subject to $A\mathbf{x}=\mathbf{b}$, note that at the minimum \mathbf{x}_0 , there can be no downhill direction consistent with the constraint $A\mathbf{x}=\mathbf{b}$. Thus, if the direction $\Delta \mathbf{x}$ at \mathbf{x}_0 is consistent with the constraint $A\mathbf{x}=\mathbf{b}$, that is $A\Delta \mathbf{x}=0$ so that $A(\mathbf{x}_0 + \Delta \mathbf{x}) = \mathbf{b}$, any subgradient ∇ for $\|\mathbf{x}\|_1$ at \mathbf{x}_0 must satisfy $\nabla^T \Delta \mathbf{x} = 0$.

A sufficient but not necessary condition for \mathbf{x}_0 to be a minimum is that there exists some \mathbf{w} such that the sub gradient at \mathbf{x}_0 is given by $\nabla = A^T \mathbf{w}$. Then for any $\Delta \mathbf{x}$ such that $A\Delta \mathbf{x} = 0$, $\nabla^T \Delta \mathbf{x} = \mathbf{w}^T A \Delta \mathbf{x} = \mathbf{w}^T \cdot \mathbf{0} = 0$. That is, for any direction consistent with the constraint $A\mathbf{x} = \mathbf{b}$, the subgradient is zero and hence \mathbf{x}_0 is a minimum.

10.3.2 The Exact Reconstruction Property

Theorem 10.3 below gives a condition that guarantees that a solution \mathbf{x}_0 to $A\mathbf{x} = \mathbf{b}$ is the unique minimum 1-norm solution to $A\mathbf{x} = \mathbf{b}$. This is a sufficient condition, but not necessary condition.

Theorem 10.3 Suppose \mathbf{x}_0 satisfies $A\mathbf{x}_0 = \mathbf{b}$. If there is a subgradient ∇ to the 1-norm function at \mathbf{x}_0 for which there exists a \mathbf{w} where $\nabla = A^T \mathbf{w}$ and the columns of A corresponding to nonzero components of \mathbf{x}_0 are linearly independent, then \mathbf{x}_0 minimizes $\|\mathbf{x}\|_1$ subject to $A\mathbf{x}=\mathbf{b}$. Furthermore, these conditions imply that \mathbf{x}_0 is the unique minimum.

Proof: We first show that \mathbf{x}_0 minimizes $\|\mathbf{x}\|_1$. Suppose \mathbf{y} is another solution to $A\mathbf{x} = \mathbf{b}$. We need to show that $||\mathbf{y}||_1 \ge ||\mathbf{x}_0||_1$. Let $\mathbf{z} = \mathbf{y} - \mathbf{x}_0$. Then $A\mathbf{z} = A\mathbf{y} - A\mathbf{x}_0 = \mathbf{0}$. Hence, $\nabla^T \mathbf{z} = (A^T \mathbf{w})^T \mathbf{z} = \mathbf{w}^T A \mathbf{z} = 0$. Now, since ∇ is a subgradient of the 1-norm function at \mathbf{x}_0 ,

 $||\mathbf{y}||_1 = ||\mathbf{x_0} + \mathbf{z}||_1 \ge ||\mathbf{x_0}||_1 + \nabla^T \cdot \mathbf{z} = ||\mathbf{x_0}||_1$

and so we have that $||\mathbf{x}_0||_1$ minimizes $||\mathbf{x}||_1$ over all solutions to $A\mathbf{x} = \mathbf{b}$.

Suppose $\tilde{\mathbf{x}}_0$ were another minimum. Then ∇ is also a subgradient at $\tilde{\mathbf{x}}_0$ as it is at \mathbf{x}_0 . To see this, for $\Delta \mathbf{x}$ such that $A\Delta \mathbf{x} = 0$,

$$\left\|\tilde{\mathbf{x}}_{0} + \Delta \mathbf{x}\right\|_{1} = \left\|\mathbf{x}_{0} + \underbrace{\tilde{\mathbf{x}}_{0} - \mathbf{x}_{0} + \Delta \mathbf{x}}_{\alpha}\right\|_{1} \ge \left\|\mathbf{x}_{0}\right\|_{1} + \nabla^{T} \left(\tilde{\mathbf{x}}_{0} - \mathbf{x}_{0} + \Delta \mathbf{x}\right).$$

The above equation follows from the definition of ∇ being a subgradient for the one norm function, $\|\|_1$, at \mathbf{x}_0 . Thus,

$$\|\tilde{\mathbf{x}}_{\mathbf{0}} + \Delta \mathbf{x}\|_{1} \ge \|\mathbf{x}_{\mathbf{0}}\|_{1} + \nabla^{T} (\tilde{\mathbf{x}}_{\mathbf{0}} - \mathbf{x}_{\mathbf{0}}) + \nabla^{T} \Delta \mathbf{x}.$$

But

$$\nabla^{T} \left(\tilde{\mathbf{x}}_{\mathbf{0}} - \mathbf{x}_{\mathbf{0}} \right) = \mathbf{w}^{T} A \left(\tilde{\mathbf{x}}_{\mathbf{0}} - \mathbf{x}_{\mathbf{0}} \right) = \mathbf{w}^{T} \left(\mathbf{b} - \mathbf{b} \right) = 0.$$

Hence, since $\tilde{\mathbf{x}_0}$ being a minimum means $||\tilde{\mathbf{x}_0}||_1 = ||\mathbf{x_0}||_1$,

$$\|\tilde{\mathbf{x}}_{\mathbf{0}} + \Delta \mathbf{x}\|_{1} \ge \|\mathbf{x}_{\mathbf{0}}\|_{1} + \nabla^{T} \Delta \mathbf{x} = \|\tilde{\mathbf{x}}_{\mathbf{0}}\|_{1} + \nabla^{T} \Delta \mathbf{x}.$$

This implies that ∇ is a sub gradient at $\tilde{\mathbf{x}}_{\mathbf{0}}$.

Now, ∇ is a subgradient at both $\mathbf{x_0}$ and $\tilde{\mathbf{x}_0}$. By Proposition 10.2, we must have that $(\nabla)_i = \operatorname{sgn}((x_0)_i) = \operatorname{sgn}((\tilde{x}_0)_i)$, whenever either is nonzero and $|(\nabla)_i| < 1$, whenever either is 0. It follows that $\mathbf{x_0}$ and $\tilde{\mathbf{x_0}}$ have the same sparseness pattern. Since $A\mathbf{x_0} = \mathbf{b}$ and $A\tilde{\mathbf{x_0}} = \mathbf{b}$ and $\mathbf{x_0}$ are both nonzero on the same coordinates, and by the assumption that the columns of A corresponding to the nonzeros of $\mathbf{x_0}$ and $\tilde{\mathbf{x_0}}$ are independent, it must be that $\mathbf{x_0} = \tilde{\mathbf{x_0}}$.

10.3.3 Restricted Isometry Property

Next we introduce the restricted isometry property that plays a key role in exact reconstruction of sparse vectors. A matrix A satisfies the *restricted isometry property*, RIP, if for any s-sparse **x** there exists a δ_s such that

$$(1 - \delta_s) |\mathbf{x}|^2 \le |A\mathbf{x}|^2 \le (1 + \delta_s) |\mathbf{x}|^2.$$
 (10.1)

Isometry is a mathematical concept; it refers to linear transformations that exactly preserve length such as rotations. If A is an $n \times n$ isometry, all its eigenvalues are ± 1 and it represents a coordinate system. Since a pair of orthogonal vectors are orthogonal in all coordinate system, for an isometry A and two orthogonal vectors \mathbf{x} and \mathbf{y} , $\mathbf{x}^T A^T A \mathbf{y} = 0$. We will prove approximate versions of these properties for matrices A satisfying the restricted isometry property. The approximate versions will be used in the sequel.

A piece of notation will be useful. For a subset S of columns of A, let A_S denote the submatrix of A consisting of the columns of S.

Lemma 10.4 If A satisfies the restricted isometry property, then

- 1. For any subset S of columns with |S| = s, the singular values of A_S are all between $1 \delta_s$ and $1 + \delta_s$.
- 2. For any two orthogonal vectors \mathbf{x} and \mathbf{y} , with supports of size s_1 and s_2 respectively, $|\mathbf{x}^T A^T A \mathbf{y}| \leq 5 |\mathbf{x}| |\mathbf{y}| (\delta_{s_1} + \delta_{s_2}).$

Proof: Item 1 follows from the definition. To prove the second item, assume without loss of generality that $|\mathbf{x}| = |\mathbf{y}| = 1$. Since \mathbf{x} and \mathbf{y} are orthogonal, $|\mathbf{x} + \mathbf{y}|^2 = 2$. Consider $|A(\mathbf{x}+\mathbf{y})|^2$. This is between $2(1-\delta_{s_1}+\delta_{s_2})^2$ and $2(1+\delta_{s_1}+\delta_{s_2})^2$ by the restricted isometry property. Also $|A\mathbf{x}|^2$ is between $(1-\delta_{s_1})^2$ and $(1+\delta_{s_1})^2$ and $|A\mathbf{y}|^2$ is between $(1-\delta_{s_2})^2$ and $(1+\delta_{s_2})^2$. Since

$$2\mathbf{x}^{T}A^{T}A\mathbf{y} = (\mathbf{x} + \mathbf{y})^{T}A^{T}A(\mathbf{x} + \mathbf{y}) - \mathbf{x}^{T}A^{T}A\mathbf{x} - \mathbf{y}^{T}A^{T}A\mathbf{y}$$
$$= |A(\mathbf{x} + \mathbf{y})|^{2} - |A\mathbf{x}|^{2} - |A\mathbf{y}|^{2},$$

it follows that

$$|2\mathbf{x}^{T}A^{T}A\mathbf{y}| \leq 2(1+\delta_{s_{1}}+\delta_{s_{2}})^{2} - (1-\delta_{s_{1}})^{2} - (1-\delta_{s_{2}})^{2}$$

$$6(\delta_{s_{1}}+\delta_{s_{2}}) + (\delta_{s_{1}}^{2}+\delta_{s_{2}}^{2}+4\delta_{s_{1}}+4\delta_{s_{2}}) \leq 9(\delta_{s_{1}}+\delta_{s_{2}}).$$

Thus, for arbitrary \mathbf{x} and $\mathbf{y} |\mathbf{x}^T A^T A \mathbf{y}| \le (9/2) |\mathbf{x}| |\mathbf{y}| (\delta_{s_1} + \delta_{s_2}).$

Theorem 10.5 Suppose A satisfies the restricted isometry property with

$$\delta_{s+1} \le \frac{1}{10\sqrt{s}}.$$

Suppose $\mathbf{x_0}$ has at most s nonzero coordinates and satisfies $A\mathbf{x} = \mathbf{b}$. Then a subgradient $\nabla ||(\mathbf{x_0})||_1$ for the 1-norm function exists at $\mathbf{x_0}$ which satisfies the conditions of Theorem 10.3 and so $\mathbf{x_0}$ is the unique minimum 1-norm solution to $A\mathbf{x} = \mathbf{b}$.

Proof: Let

$$S = \{i | (\mathbf{x_0})_i \neq 0\}$$

be the support of $\mathbf{x_0}$ and let $\overline{S} = \{i | (\mathbf{x_0})_i = \mathbf{0}\}$ be the complement set of coordinates. To find a subgradient \mathbf{u} at $\mathbf{x_0}$ satisfying Theorem 10.3, search for a \mathbf{w} such that $\mathbf{u} = A^T \mathbf{w}$ where for coordinates in which $\mathbf{x_0} \neq 0$, $\mathbf{u} = sgn(\mathbf{x_0})$ and for the remaining coordinates the 2-norm of \mathbf{u} is minimized. Solving for \mathbf{w} is a least squares problem. Let \mathbf{z} be the vector with support S, with $z_i = sgn(\mathbf{x_0})$ on S. Consider the vector \mathbf{w} defined by

$$\mathbf{w} = A_S \left(A_S^T A_S \right)^{-1} \mathbf{z}$$

This happens to be the solution of the least squares problem, but we do not need this fact. We only state it to tell the reader how we came up with this expression. Note that A_S has independent columns from the restricted isometry property assumption, and so

 $A_S^T A_S$ is invertible. We will prove that this **w** satisfies the conditions of Theorem 10.3. First, for coordinates in S,

$$(A^T \mathbf{w})_S = (A_S)^T A_S (A_S^T A_S)^{-1} \mathbf{z} = \mathbf{z}$$

as required.

For coordinates in \overline{S} , we have

$$(A^T \mathbf{w})_{\bar{S}} = (A_{\bar{S}})^T A_S (A_S^T A_S)^{-1} \mathbf{z}.$$

Now, the eigenvalues of $A_S^T A_S$, which are the squares of the singular values of A_S , are between $(1 - \delta_s)^2$ and $(1 + \delta_s)^2$. So $||(A_S^T A_S)^{-1}|| \leq \frac{1}{(1 - \delta_S)^2}$. Letting $\mathbf{p} = (A_S^T A_S)^{-1} \mathbf{z}$, we have $|\mathbf{p}| \leq \frac{\sqrt{s}}{(1 - \delta_S)^2}$. Write $A_s \mathbf{p}$ as $A \mathbf{q}$, where \mathbf{q} has all coordinates in \overline{S} equal to zero. Now, for $j \in \overline{S}$

$$(A^T \mathbf{w})_j = e_j^T A^T A \mathbf{q}$$

and part (2) of Lemma 10.4 gives $|(A^T \mathbf{w})_j| \leq 9\delta_{s+1}\sqrt{s}/(1-\delta_s^2) \leq 1/2$ establishing the Theorem 10.3 holds.

A Gaussian matrix is a matrix where each element is an independent Gaussian variable. Gaussian matrices satisfy the restricted isometry property. (Exercise ??)

10.4 Applications

10.4.1 Sparse Vector in Some Coordinate Basis

Consider $A\mathbf{x} = \mathbf{b}$ where A is a square $n \times n$ matrix. The vectors \mathbf{x} and \mathbf{b} can be considered as two representations of the same quantity. For example, \mathbf{x} might be a discrete time sequence with \mathbf{b} the frequency spectrum of \mathbf{x} and the matrix A the Fourier transform. The quantity \mathbf{x} can be represented in the time domain by \mathbf{x} and in the frequency domain by its Fourier transform \mathbf{b} . In fact, any orthonormal matrix can be thought of as a transformation and there are many important transformations other than the Fourier transformation.

Consider a transformation A and a signal \mathbf{x} in some standard representation. Then $\mathbf{y} = A\mathbf{x}$ transforms the signal \mathbf{x} to another representation \mathbf{y} . If A spreads any sparse signal \mathbf{x} out so that the information contained in each coordinate in the standard basis is spread out to all coordinates in the second basis, then the two representations are said to be *incoherent*. A signal and its Fourier transform are one example of incoherent vectors. This suggests that if \mathbf{x} is sparse, only a few randomly selected coordinates of its Fourier transform are needed to reconstruct \mathbf{x} . In the next section we show that a signal cannot be too sparse in both its time domain and its frequency domain.

10.4.2 A Representation Cannot be Sparse in Both Time and Frequency Domains

We now show that there is an uncertainty principle that states that a time signal cannot be sparse in both the time domain and the frequency domain. If the signal is of length n, then the product of the number of nonzero coordinates in the time domain and the number of nonzero coordinates in the frequency domain must be at least n. We first prove two technical lemmas.

In dealing with the Fourier transform it is convenient for indices to run from 0 to n-1 rather than from 1 to n. Let $x_0, x_1, \ldots, x_{n-1}$ be a sequence and let $f_0, f_1, \ldots, f_{n-1}$ be its discrete Fourier transform. Let $i = \sqrt{-1}$. Then $f_j = \frac{1}{\sqrt{n}} \sum_{k=0}^{n-1} x_k e^{-\frac{2\pi i}{n}jk}$, $j = 0, \ldots, n-1$. In matrix form $\mathbf{f} = Z\mathbf{x}$ where $z_{jk} = e^{-\frac{2\pi i}{n}jk}$.

$$\begin{pmatrix} f_0 \\ f_1 \\ \vdots \\ f_{n-1} \end{pmatrix} = \frac{1}{\sqrt{n}} \begin{pmatrix} 1 & 1 & 1 & \cdots & 1 \\ 1 & e^{-\frac{2\pi i}{n}} & e^{-\frac{2\pi i}{n}2} & \cdots & e^{-\frac{2\pi i}{n}(n-1)} \\ \vdots & \vdots & & \vdots \\ 1 & e^{-\frac{2\pi i}{n}(n-1)} & e^{-\frac{2\pi i}{n}2(n-1)} & \cdots & e^{-\frac{2\pi i}{n}(n-1)^2} \end{pmatrix} \begin{pmatrix} x_0 \\ x_1 \\ \vdots \\ x_{n-1} \end{pmatrix}$$

If some of the elements of \mathbf{x} are zero, delete the zero elements of \mathbf{x} and the corresponding columns of the matrix. To maintain a square matrix, let n_x be the number of nonzero elements in \mathbf{x} and select n_x consecutive rows of the matrix. Normalize the columns of the resulting submatrix by dividing each element in a column by the column element in the first row. The resulting submatrix is a Vandermonde matrix that looks like

$$\left(\begin{array}{rrrrr} 1 & 1 & 1 & 1 \\ a & b & c & d \\ a^2 & b^2 & c^2 & d^2 \\ a^3 & b^3 & c^3 & d^3 \end{array}\right)$$

and is nonsingular.

Lemma 10.6 If $x_0, x_1, \ldots, x_{n-1}$ has n_x nonzero elements, then $f_0, f_1, \ldots, f_{n-1}$ cannot have n_x consecutive zeros.

Proof: Let $i_1, i_2, \ldots, i_{n_x}$ be the indices of the nonzero elements of **x**. Then the elements of the Fourier transform in the range $k = m + 1, m + 2, \ldots, m + n_x$ are

$$f_k = \frac{1}{\sqrt{n}} \sum_{j=1}^{n_x} x_{i_j} e^{\frac{-2\pi i}{n} k i_j}$$

Note the use of i as $\sqrt{-1}$ and the multiplication of the exponent by i_j to account for the actual location of the element in the sequence. Normally, if every element in the sequence

was included, we would just multiply by the index of summation.

Convert the equation to matrix form by defining $z_{kj} = \frac{1}{\sqrt{n}} \exp(-\frac{2\pi i}{n} k i_j)$ and write $\mathbf{f} = Z\mathbf{x}$. Actually instead of \mathbf{x} , write the vector consisting of the nonzero elements of \mathbf{x} . By its definition, $\mathbf{x} \neq 0$. To prove the lemma we need to show that \mathbf{f} is nonzero. This will be true provided Z is nonsingular. If we rescale Z by dividing each column by its leading entry we get the Vandermonde determinant which is nonsingular.

Theorem 10.7 Let n_x be the number of nonzero elements in \mathbf{x} and let n_f be the number of nonzero elements in the Fourier transform of \mathbf{x} . Let n_x divide n. Then $n_x n_f \ge n$.

Proof: If **x** has n_x nonzero elements, **f** cannot have a consecutive block of n_x zeros. Since n_x divides n there are $\frac{n}{n_x}$ blocks each containing at least one nonzero element. Thus, the product of nonzero elements in **x** and **f** is at least n.

Fourier transform of spikes prove that above bound is tight

To show that the bound in Theorem 10.7 is tight we show that the Fourier transform of the sequence of length n consisting of \sqrt{n} ones, each one separated by $\sqrt{n} - 1$ zeros, is the sequence itself. For example, the Fourier transform of the sequence 100100100 is 100100100. Thus, for this class of sequences, $n_x n_f = n$.

Theorem 10.8 Let $S(\sqrt{n}, \sqrt{n})$ be the sequence of 1's and 0's with \sqrt{n} 1's spaced \sqrt{n} apart. The Fourier transform of $S(\sqrt{n}, \sqrt{n})$ is itself.

Proof: Consider the columns $0, \sqrt{n}, 2\sqrt{n}, \ldots, (\sqrt{n}-1)\sqrt{n}$. These are the columns for which $S(\sqrt{n}, \sqrt{n})$ has value 1. The element of the matrix Z in the row $j\sqrt{n}$ of column $k\sqrt{n}, 0 \le k < \sqrt{n}$ is $z^{nkj} = 1$. Thus, for these rows Z times the vector $S(\sqrt{n}, \sqrt{n}) = \sqrt{n}$ and the $1/\sqrt{n}$ normalization yields $f_{j\sqrt{n}} = 1$.

For rows whose index is not of the form $j\sqrt{n}$, the row $b, b \neq j\sqrt{n}, j \in \{0, \sqrt{n}, \dots, \sqrt{n-1}\}$, the elements in row b in the columns $0, \sqrt{n}, 2\sqrt{n}, \dots, (\sqrt{n-1})\sqrt{n}$ are $1, z^b, z^{2b}, \dots, z^{(\sqrt{n-1})b}$ and thus $f_b = \frac{1}{\sqrt{n}} \left(1 + z^b + z^{2b} \dots + z^{(\sqrt{n-1})b}\right) = \frac{1}{\sqrt{n}} \frac{z^{\sqrt{n}b} - 1}{z^{-1}} = 0$ since $z^{b\sqrt{n}} = 1$ and $z \neq 1$.

Uniqueness of l_1 optimization

Consider a redundant representation for a sequence. One such representation would be representing a sequence as the concatenation of two sequences, one specified by its coordinates and the other by its Fourier transform. Suppose some sequence could be represented as a sequence of coordinates and Fourier coefficients sparsely in two different ways. Then by subtraction, the zero sequence could be represented by a sparse sequence. The representation of the zero sequence cannot be solely coordinates or Fourier coefficients. If y is the coordinate sequence in the representation of the zero sequence, then the Fourier portion of the representation must represent -y. Thus y and its Fourier transform would have sparse representations contradicting $n_x n_f \geq n$. Notice that a factor of two comes in

1	1	1	1	1	1	1	1	1
1	z	z^2	z^3	z^4	z^5	z^6	z^7	z^8
1	z^2	z^4	z^6	z^8	z	z^3	z^5	z^7
1	z^3	z^6	1	z^3	z^6	1	z^3	z^6
1	z^4	z^8	z^3	z^7	z^2	z^6	z	z^5
1	z^5	z	z^6	z^2	z^7	z^3	z^8	z^4
1	z^6	z^3	1	z^6	z^3	1	z^6	z^3
1	z^7	z^5	z^3	z	z^8	z^6	z^4	z^2
1	z^8	z^7	z^6	z^5	z^4	z^3	z^2	z

Figure 10.5: The matrix Z for n=9.

when we subtract the two representations.

Suppose two sparse signals had Fourier transforms that agreed in almost all of their coordinates. Then the difference would be a sparse signal with a sparse transform. This is not possible. Thus, if one selects $\log n$ elements of their transform these elements should distinguish between these two signals.

10.4.3 Biological

There are many areas where linear systems arise in which a sparse solution is unique. One is in plant breading. Consider a breeder who has a number of apple trees and for each tree observes the strength of some desirable feature. He wishes to determine which genes are responsible for the feature so he can cross bread to obtain a tree that better expresses the desirable feature. This gives rise to a set of equations $A\mathbf{x} = \mathbf{b}$ where each row of the matrix A corresponds to a tree and each column to a position on the genone. See Figure 10.6. The vector \mathbf{b} corresponds to the strength of the desired feature in each tree. The solution \mathbf{x} tells us the position on the genone corresponding to the genes that account for the feature. It would be surprising if there were two small independent sets of genes that accounted for the desired feature. Thus, the matrix must have a property that allows only one sparse solution.

10.4.4 Finding Overlapping Cliques or Communities

Consider a graph that consists of several cliques. Suppose we can observe only low level information such as edges and we wish to identify the cliques. An instance of this problem is the task of identifying which of ten players belongs to which of two teams of five players each when one can only observe interactions between pairs of individuals. There is an interaction between two players if and only if they are on the same team. In this situation we have a matrix A with $\binom{10}{5}$ columns and $\binom{10}{2}$ rows. The columns represent possible teams and the rows represent pairs of individuals. Let **b** be the $\binom{10}{2}$ dimensional vector of observed interactions. Let **x** be a solution to $A\mathbf{x} = \mathbf{b}$. There is a

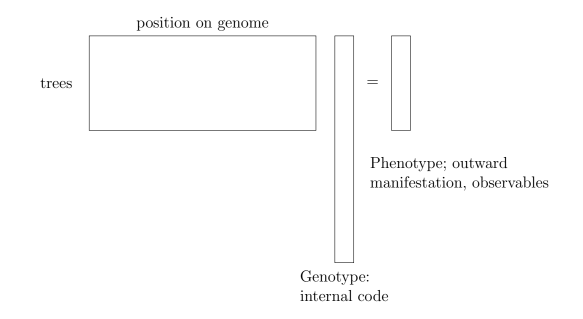


Figure 10.6: The system of linear equations used to find the internal code for some observable phenomenon.

sparse solution \mathbf{x} where \mathbf{x} is all zeros except for the two 1's for 12345 and 678910 where the two teams are $\{1,2,3,4,5\}$ and $\{6,7,8,9,10\}$. The question is can we recover \mathbf{x} from \mathbf{b} . If the matrix A had satisfied the restricted isometry condition, then we could surely do this. Although A does not satisfy the restricted isometry condition which guarantees recover of all sparse vectors, we can recover the sparse vector in the case where the teams are non overlapping or almost non overlapping. If A satisfied the restricted isometry property we would minimize $\|\mathbf{x}\|_1$ subject to $A\mathbf{x} = \mathbf{b}$. Instead, we minimize $\|\mathbf{x}\|_1$ subject to $\|A\mathbf{x} - \mathbf{b}\|_{\infty} \leq \varepsilon$ where we bound the largest error.

10.4.5 Low Rank Matrices

Suppose L is a low rank matrix that has been corrupted by noise. That is, M = L + R. If the R is Gaussian, then principle component analysis will recover L from M. However, if L has been corrupted by several missing entries or several entries have a large noise added to them and they become outliers, then principle component analysis may be far off. However, if L is low rank and R is sparse, then L can be recovered effectively from L+R. To do this, find the L and R that minimize $||L||_F^2 + \lambda ||R||_1$. Here $||L||_F^2$ is the sum of the singular values of L. A small value of $||L||_F^2$ indicates a low rank matrix. Notice that we do not need to know the rank of L or the elements that were corrupted. All we need is that the low rank matrix L is not sparse and that the sparse matrix R is not low rank. We leave the proof as an exercise.

An example where low rank matrices that have been corrupted might occur is aerial

photographs of an intersection. Given a long sequence of such photographs, they will be the same except for cars and people. If each photo is converted to a vector and the vector used to make a column of a matrix, then the matrix will be low rank corrupted by the traffic. Finding the original low rank matrix will separate the cars and people from the back ground.

10.5 Gradient

The gradient of a function $f(\mathbf{x})$ of d variables, $\mathbf{x} = (x_1, x_2, \dots, x_d)$, at a point \mathbf{x}_0 is denoted $\nabla f(\mathbf{x}_0)$. It is a d-dimensional vector with components $\frac{\partial f}{\partial x_1}(\mathbf{x}_0), \frac{\partial f}{\partial x_2}(\mathbf{x}_0), \dots, \frac{\partial f}{\partial x_d}(\mathbf{x}_0)$, where $\frac{\partial f}{\partial x_i}$ are partial derivatives. Without explicitly stating, we assume that the derivatives referred to exist. The rate of increase of the function f as we move from \mathbf{x}_0 in a direction **u** is easily seen to be $\nabla f(\mathbf{x}_0) \cdot \mathbf{u}$. So the direction of steepest descent is $-\nabla f(\mathbf{x_0})$; this is a natural direction to move in if we wish to minimize f. But by how much should we move? A large move may overshoot the minimum. [See figure (10.7).] A simple fix is to minimize f on the line from \mathbf{x}_0 in the direction of steepest descent by solving a one dimensional minimization problem. This gets us the next iterate $\mathbf{x_1}$ and we may repeat. Here, we will not discuss the issue of step-size any further. Instead, we focus on "infinitesimal" gradient descent, where, the algorithm makes infinitesimal moves in the $-\nabla f(\mathbf{x}_0)$ direction. Whenever ∇f is not the zero vector, we strictly decrease the function in the direction $-\nabla \mathbf{f}$, so the current point is not a minimum of the function f. Conversely, a point **x** where $\nabla \mathbf{f} = \mathbf{0}$ is called a *first-order local optimum* of f. In general, local minima do not have to be global minima (see (10.7)) and gradient descent may converge to a local minimum which is not a global minimum. In the special case when the function f is convex, this is not the case. A function f of a single variable x is said to be convex if for any two points x and y, the line joining f(x) and f(y) is above the curve $f(\cdot)$. A function of many variables is convex if on any line segment in its domain, it acts as a convex function of one variable on the line segment.

Definition 10.1 A function f over a convex domain is a convex function if for any two points \mathbf{x}, \mathbf{y} in the domain, and any λ in [0, 1] we have

$$f(\lambda \mathbf{x} + (1 - \lambda)\mathbf{y}) \le \lambda f(\mathbf{x}) + (1 - \lambda)f(\mathbf{y}).$$

The function is concave if the inequality is satisfied with \geq instead of \leq .

Theorem 10.9 Suppose f is a convex, differentiable, function defined on a closed bounded convex domain. Then any first-order local minimum is also a global minimum. Thus, infinitesimal gradient descent always reaches the global minimum.

Proof: We will prove that if \mathbf{x} is a local minimum, then it must be a global minimum. If not, consider a global minimum point $\mathbf{y} \neq \mathbf{x}$. But on the line joining \mathbf{x} and \mathbf{y} , the function must not go above the line joining $f(\mathbf{x})$ and $f(\mathbf{y})$. This means for an infinitesimal $\varepsilon > 0$, if we move distance ε from \mathbf{x} towards \mathbf{y} , the function must decrease, so we cannot have $\nabla \mathbf{f}(\mathbf{x}) = \mathbf{0}$, contradicting the assumption that \mathbf{x} is a local minimum.

The second derivatives $\frac{\partial^2}{\partial x_i \partial x_j}$ form a matrix, called the Hessian, denoted $H(f(\mathbf{x}))$. The Hessian of f at \mathbf{x} is a symmetric $d \times d$ matrix with $(i, j)^{th}$ entry $\frac{\partial^2 f}{\partial x_i \partial x_j}(\mathbf{x})$. The second derivative of f at \mathbf{x} in the direction \mathbf{u} is the rate of change of the first derivative as we move along \mathbf{u} from \mathbf{x} . It is easy to see that it equals

$$\mathbf{u}^T H(f(\mathbf{x}))\mathbf{u}$$

To see this, note that the second derivative of f along **u** is

$$\sum_{j} u_{j} \frac{\partial}{\partial x_{j}} \left(\nabla f(\mathbf{x}) \cdot \mathbf{u} \right) = \sum_{j} u_{j} \sum_{i} \frac{\partial}{\partial x_{j}} \left(u_{i} \frac{\partial f}{\partial x_{i}} \right)$$
$$= \sum_{j,i} u_{j} u_{i} \frac{\partial^{2} f}{\partial x_{j} \partial x_{i}} (\mathbf{x}).$$

Theorem 10.10 Suppose f is a function from a closed convex domain D in \mathbb{R}^d to the reals and the Hessian of f exists everywhere in D. Then f is convex (concave) on D if and only if the Hessian of f is positive (negative) semi-definite everywhere on D.

Gradient descent requires the gradient to exist. But, even if the gradient is not always defined, one can minimize a convex function over a convex domain efficiently, i.e., in polynomial time. Here, the quote is added because of the lack of rigor in the statement, one can only find an approximate minimum and the time really depends on the error parameter as well as the presentation of the convex set. We do not go into these details here. But, in principle we can minimize a convex function over a convex domain. We can also maximize a concave function over a convex domain. However, in general, we do not have efficient procedures to maximize a convex function over a convex set. It is easy to see that at a first-order local minimum of a possibly non-convex function, the gradient vanishes. But second-order local decrease of the function may be possible. The steepest second-order decrease is in the direction of $\pm \mathbf{v}$, where, \mathbf{v} is the eigenvector of the Hessian corresponding to the largest absolute valued eigenvalue.

10.6 Linear Programming

Linear programming is an optimization problem which has been carefully studied and is immensely useful. We consider linear programming problem in the following form where A is an $m \times n$ matrix of rank m, c is $1 \times n$, b is $m \times 1$ and x is $n \times 1$):

max
$$\mathbf{c} \cdot \mathbf{x}$$
 subject to $A\mathbf{x} = \mathbf{b}, x \ge 0$.

Inequality constraints can be converted to this form by adding slack variables. Also, we can do Gaussian elimination on A and if it does not have rank m, we either find that the system of equations has no solution, whence we may stop or we can find and discard redundant equations. After this preprocessing, we may assume that A 's rows are independent.

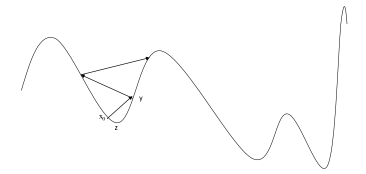


Figure 10.7

The simplex algorithm is a classical method to solve linear programming problems. It is a vast subject and is well discussed in many texts. Here, we will discuss the ellipsoid algorithm which is in a sense based more on continuous mathematics and is closer to the spirit of this book.

10.6.1 The Ellipsoid Algorithm

The first polynomial time algorithm for Linear Programming was developed by Khachian based on work of Iudin, Nemirovsky and Shor and is called the ellipsoid algorithm. The algorithm is best stated for the seemingly simpler problem of determining whether there is a solution to $A\mathbf{x} \leq \mathbf{b}$ and if so finding one. The ellipsoid algorithm starts with a large ball in *d*-space which is guaranteed to contain the polyhedron $A\mathbf{x} \leq \mathbf{b}$. Even though we do not yet know if the polyhedron is empty or non-empty, such a ball can be found. It checks if the center of the ball is in the polyhedron, if it is, we have achieved our objective. If not, we know from convex geometry (in particular, the Separating Hyperplane Theorem) that there is a hyperplane called the separating hyperplane through the center of the ball such that the whole polytope lies in one half space.

We then find an ellipsoid which contains the ball intersected with this half-space See Figure (10.8. The ellipsoid is guaranteed to contain $Ax \leq b$ as was the ball earlier. We now check if the center of the ellipsoid satisfies the inequalities. If not, there is a separating hyper plane again and we may repeat the process. After a suitable number of steps, either we find a solution to the original $A\mathbf{x} \leq \mathbf{b}$ or, we end up with a very small ellipsoid. Now if the original A and \mathbf{b} had integer entries, one can ensure that the set $A\mathbf{x} \leq \mathbf{b}$, after a slight perturbation which preserves its emptiness/non-emptiness, has a volume of at least some $\epsilon > 0$ and if our ellipsoid has shrunk to a volume of less than this ϵ , then we know there is no solution and we can stop. Clearly this must happen within $\log_{\rho} V_0/\epsilon = O((V_0 d)/\epsilon)$, where V_0 is an upper bound on the initial volume and ρ is the factor by which the volume shrinks in each step. We do not go into details of how to get a value for V_0 here, but the important points are that (i) V_0 only occurs under the logarithm and (ii) the dependence on d is linear. These features ensure a polynomial time algorithm.

The main difficulty in proving fast convergence is to show that the volume of the ellipsoid shrinks by a certain factor in each step. Thus, the question can be phrased as suppose E is an ellipsoid with center \mathbf{x}_0 and consider the half-ellipsoid E' defined by

$$E' = \{ \mathbf{x} : \mathbf{x} \in E ; \ \mathbf{a} \cdot (\mathbf{x} - \mathbf{x_0}) \ge 0 \},\$$

where, **a** is some unit length vector. Let \hat{E} be the smallest volume ellipsoid containing E'. Show that

$$\frac{\operatorname{Vol}(E)}{\operatorname{Vol}(E)} \le 1 - \rho$$

for some $\rho > 0$. A sequence of geometric reductions transforms this into a simple problem. First, observe that we can translate the entire picture and assume that $\mathbf{x}_0 = 0$.

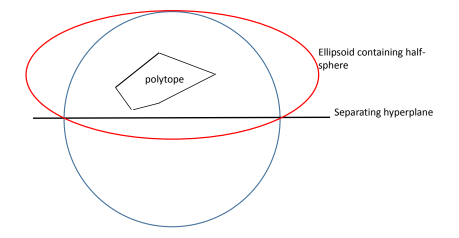


Figure 10.8: Ellipsoid Algorithm

Next, rotate the coordinate axes so that **a** is replaced by (1, 0, 0, ..., 0). Finally, make a nonsingular linear transformation τ so that $\tau E = B = \{\mathbf{x} : |\mathbf{x}| = 1\}$, the unit sphere. The important point is that a nonsingular linear transformation τ multiplies the volumes of all sets by $|\det(\tau)|$, so that $\frac{\operatorname{Vol}(\hat{E})}{\operatorname{Vol}(E)} = \frac{\operatorname{Vol}(\tau(\hat{E}))}{\operatorname{Vol}(\tau(E))}$. Now, the following lemma answers the question raised.

Lemma 10.11 Consider the half-sphere $B' = {\mathbf{x} : x_1 \ge 0 ; |\mathbf{x}| \le 1}$. The following ellipsoid \hat{E} contains B':

$$\hat{E} = \left\{ \mathbf{x} \left| \left(\frac{d+1}{d} \right)^2 \left(x_1 - \frac{1}{d+1} \right)^2 + \left(\frac{d^2 - 1}{d^2} \right) \left(x_2^2 + x_3^2 + \dots + x_d^2 \right) \le 1 \right\}.$$

Further,

$$\frac{\operatorname{Vol}(\hat{E})}{\operatorname{Vol}(B)} = \left(\frac{d}{d+1}\right) \left(\frac{d^2}{d^2-1}\right)^{(d-1)/2} \le 1 - \frac{1}{4d}$$

Proof: See Exercise (10.24).

10.7 Integer Optimization

The problem of maximizing a linear function subject to linear inequality constraints, but with the variables constrained to be integers is called integer programming:

Max $\mathbf{c} \cdot \mathbf{x}$ subject to $A\mathbf{x} \leq \mathbf{b} \ x_i$ integers.

This problem is NP-hard. One way to handle the hardness is to relax the integer constraints, solve the linear program in polynomial time and round the fractional values to integers. The simplest rounding, round each variable which is 1/2 or more to 1, the rest to 0, yields sensible results in some cases. The vertex cover problem is one of them. The problem is to choose a subset of vertices so that each edge is covered, at least one of its end points is in the subset. The integer program is:

$$\operatorname{Min}\sum_{i} x_{i} \text{ subject to } x_{i} + x_{j} \geq 1 \forall \text{ edges } (i, j); \ x_{i} \text{ integers } .$$

Solve the linear program. At least one variable for each edge must be at least 1/2 and the simple rounding converts it to 1. So the integer solution is still feasible. It clearly at most doubles the objective function from the linear programming solution and since the LP solution value is at most the optimal integer programming solution value, we get within a factor of 2 of the optimal.

10.8 Semi-Definite Programming

Semi-definite programs are special cases of convex programs. Recall that an $n \times n$ matrix A is positive semi-definite if and only if A is symmetric and for all $\mathbf{x} \in \mathbf{R}^n$, $\mathbf{x}^T A \mathbf{x} \ge 0$. There are many equivalent characterizations of positive semi-definite matrices. We mention one. A symmetric matrix A is positive semi-definite if and only if it can be expressed as $A = BB^T$ for a possibly rectangular matrix B.

A semi-definite program (SDP) is the problem of minimizing a linear function $\mathbf{c}^T \mathbf{x}$ subject to a constraint that $F = F_0 + F_1 x_1 + F_2 x_2 + \cdots + F_d x_d$ is positive semi-definite. Here F_0, F_1, \ldots, F_d are given symmetric matrices.

This is a convex program since the set of \mathbf{x} satisfying the constraint is a convex set. To see this, note that if $F(\mathbf{x}) = F_0 + F_1x_1 + F_2x_2 + \cdots + F_dx_d$ and $F(\mathbf{y}) = F_0 + F_1y_1 + F_2y_2 + \cdots + F_dy_d$ are positive semi-definite, then so is $F(\alpha \mathbf{x} + (1 - \alpha)\mathbf{y})$ for all $\alpha \in [0, 1]$. In principle, SDP's can be solved in polynomial time. It turns out that there are more efficient algorithms for SDP's than general convex programs and that many interesting problems can be formulated as SDP's. We discuss the latter aspect here.

Linear programs are special cases of SDP's. For any vector \mathbf{v} , let diag(\mathbf{v}) denote a diagonal matrix with the components of \mathbf{v} on the diagonal. Then it is easy to see that the constraints $\mathbf{v} \geq \mathbf{0}$ are equivalent to the constraint diag(\mathbf{v}) is positive semi-definite. Consider the linear program:

Minimize $\mathbf{c}^T \mathbf{x}$ subject to $A\mathbf{x} = \mathbf{b}$; $\mathbf{x} \ge \mathbf{0}$. Rewrite $A\mathbf{x} = \mathbf{b}$ as $A\mathbf{x} - \mathbf{b} \ge \mathbf{0}$; $\mathbf{b} - A\mathbf{x} \ge \mathbf{0}$ and use the idea of diagonal matrices above to formulate this as an SDP.

A second interesting example is that of quadratic programs of the form:

xc

Minimize
$$\frac{(\mathbf{c}^T \mathbf{x})^2}{\mathbf{d}^T \mathbf{x}}$$
 subject to $A\mathbf{x} + \mathbf{b} \ge \mathbf{0}$.

This is equivalent to

Minimize t subject to $A\mathbf{x} + \mathbf{b} \ge \mathbf{0}$ and $t \ge \frac{(\mathbf{c}^T \mathbf{x})^2}{\mathbf{d}^T \mathbf{x}}$.

This is in turn equivalent to the SDP

Minimize t subject to the following matrix being positive semi-definite:

$$\left(\begin{array}{ccc} \operatorname{diag}(A\mathbf{x} + \mathbf{b}) & 0 & 0\\ 0 & t & \mathbf{c}^{T}\mathbf{x}\\ 0 & \mathbf{c}^{T}\mathbf{x} & \mathbf{d}^{T}\mathbf{x} \end{array}\right)$$

An exciting area of application of SDP is to solve some integer problems. The central idea is best illustrated by its early application in a breakthrough due to Goemans and Williamson ([?]) for the maximum cut problem which given a graph G(V, E) asks for the cut S, \overline{S} maximizing the number of edges going across the cut from S to \overline{S} . For each $i \in V$, let x_i be an integer variable assuming values ± 1 depending on whether $i \in S$ or $i \in \overline{S}$ respectively. Then the max-cut problem can be posed as

Maximize $\sum_{(i,j)\in E} (1-x_i x_j)$ subject to the constraints $x_i \in \{-1,+1\}$.

The integrality constraint on the x_i makes the problem NP-hard. Instead replace the integer constraints by allowing the \mathbf{x}_i to be unit length vectors. This enlarges the set of feasible solutions since ± 1 are just 1-dimensional vectors of length 1. The relaxed problem is an SDP and can be solved in polynomial time. To see that it is an SDP, consider \mathbf{x}_i as the rows of a matrix X. The variables of our SDP are not X, but actually $Y = XX^T$, which is a positive definite matrix. The SDP is

Maximize $\sum_{(i,j)\in E} (1-y_{ij})$ subject to Y positive semi-define

which can be solved in polynomial time. From the solution Y, find X satisfying $Y = XX^T$. Now, instead of a ± 1 label on each vertex, we have vector labels, namely the rows of X. We need to round the vectors to ± 1 to get an S. One natural way to do this is to pick a random vector \mathbf{v} and if for vertex i, $\mathbf{x}_i \cdot \mathbf{v}$ is positive, put i in S, otherwise put it in \overline{S} . Goemans and Wiiliamson showed that this method produces a cut guaranteed to be at least 0.878 times the maximum. The .878 factor is a big improvement on the previous best factor of 0.5 which is easy to get by putting each vertex into S with probability 1/2.

10.9 Exercises

Exercise 10.1 Select a method that you believe is good for combining individual rankings into a global ranking. Consider a set of rankings where each individual ranks b last. One by one move b from the bottom to the top leaving the other rankings in place. Does there exist a v_b as in Theorem 10.1 where v_b is the ranking that causes b to move from the bottom to the top in the global ranking. If not, does your method of combing individual rankings rankings satisfy the axioms of unanimity and independence of irrelevant alternatives.

Exercise 10.2 Show that the three axioms: non dictator, unanimity, and independence of irrelevant alternatives are independent.

Exercise 10.3 Does the axiom of independence of irrelevant alternatives make sense? What if there were three rankings of five items. In the first two rankings, A is number one and B is number two. In the third ranking, B is number one and A is number five. One might compute an average score where a low score is good. A gets a score of 1+1+5=7 and B gets a score of 2+2+1=5 and B is ranked number one in the global raking. Now if the third ranker moves A up to the second position, A's score becomes 1+1+2=4 and the global ranking of A and B changes even though no individual ranking of A and B changed. Is there some alternative axiom to replace independence of irrelevant alternatives? Write a paragraph on your thoughts on this issue.

Exercise 10.4 Prove that the global ranking agrees with column v_b even if b is moved down through the column.

Exercise 10.5 Create a random 100 by 100 orthonormal matrix A and a sparse 100dimensional vector \mathbf{x} . Compute $A\mathbf{x} = \mathbf{b}$. Randomly select a few coordinates of \mathbf{b} and reconstruct \mathbf{x} from the samples of \mathbf{b} using the minimization of 1-norm technique of Section 10.3.1. Did you get \mathbf{x} back?

Exercise 10.6 Let A be a low rank $n \times m$ matrix. Let r be the rank of A. Let \tilde{A} be A corrupted by Gaussian noise. Prove that the rank r SVD approximation to \tilde{A} minimizes $\left|A - \tilde{A}\right|_{F}^{2}$.

Exercise 10.7 Prove that minimizing $||x||_0$ subject to Ax = b is NP-complete.

Exercise 10.8 Let A be a Gaussian matrix where each element is a random Gauussian variable with zero mean and variance one. Prove that A has the restricted isometry property.

Exercise 10.9 Generate 100×100 matrices of rank 20, 40, 60 80, and 100. In each matrix randomly delete 50, 100, 200, or 400 entries. In each case try to recover the original matrix. How well do you do?

Exercise 10.10 Repeat the previous exercise but instead of deleting elements, corrupt the elements by adding a reasonable size corruption to the randomly selected matrix entires.

Exercise 10.11 Compute the Fourier transform of the sequence 1000010000.

Exercise 10.12 What is the Fourier transform of a Gaussian?

Exercise 10.13 What is the Fourier transform of a cyclic shift?

Exercise 10.14 Let S(i, j) be the sequence of *i* blocks each of length *j* where each block of symbols is a 1 followed by i - 1 0's. The number n=6 is factorable but not a perfect square. What is Fourier transform of S(2,3) = 100100?

Exercise 10.15 Let Z be the n root of unity. Prove that $\{z^{bi}|0 \le i < n\} = \{z^i|0 \le i < n\}$ provide that b does not divide n.

 $\left(\begin{array}{ccccc} 1 & 1 & \cdots & 1 \\ a & b & \cdots & c \\ a^2 & b^2 & \cdots & c^2 \\ \vdots & \vdots & & \vdots \\ a^d & b^d & \cdots & c^d \end{array}\right)$

Show that if the elements in the second row of the Vandermonde matrix are distinct, then the Vandermonde matrix is nonsingular by using the fact that specifying the value of an n^{th} degree polynomial at n + 1 points uniquely determines the polynomial.

Exercise 10.16 Many problems can be formulated as finding \mathbf{x} satisfying $A\mathbf{x} = \mathbf{b} + \mathbf{r}$ where \mathbf{r} is some residual error. Discuss the advantages and disadvantages of each of the following three versions of the problem.

- 1. Set $\mathbf{r} = 0$ and find $\mathbf{x} = argmin \|\mathbf{x}\|_1$ satisfying $A\mathbf{x} = \mathbf{b}$
- 2. Lasso: find $\mathbf{x} = argmin\left(\|\mathbf{x}\|_1 + \alpha \|\mathbf{r}\|_2^2\right)$ satisfying $A\mathbf{x} = \mathbf{b}$
- 3. find $\underline{x} = argmin \|\mathbf{x}\|_1$ such that $\|\mathbf{r}\|_2 < \varepsilon$

Exercise 10.17 Create a graph of overlapping communities as follows. Let n=1,000. Partition the integers into ten blocks each of size 100. The first block is $\{1, 2, ..., 100\}$. The second is $\{100, 101, ..., 200\}$, and so on. Add edges to the graph so that the vertices in each block form a clique. Now randomly permute the indices and partition the sequence into ten blocks of 100 vertices each. Again add edges so that these new blocks are cliques. Randomly permute the indices a second time and repeat the process of adding edges. The result is a graph in which each vertex is in three cliques. Explain how to find the cliques given the graph.

Exercise 10.18 Repeat the above exercise but instead of adding edges to form cliques, use each block to form a G(100,p) graph. For how small a p can you recover the blocks? What if you add G(1,000,q) to the graph for some small value of q.

Exercise 10.19 Construct an $n \times m$ matrix A where each of the m columns is a 0-1 indicator vector with approximately 1/4 entries being 1. Then $B = AA^T$ is a symmetric matrix that can be viewed as the adjacency matrix of an n vertex graph. Some edges will have weight greater than one. The graph consists of a number of possibly over lapping cliques. Your task given B is to find the cliques by the following technique of finding a 0-1 vector in the column space of B by the following linear program for finding b and x.

$$b = argmin||b||_1$$

subject to

$$\begin{aligned} Bx &= b\\ b_1 &= 1\\ 0 &\leq b_i \leq 1 \quad 2 \leq i \leq n \end{aligned}$$

Then subtract bb^T from B and repeat.

Exercise 10.20 Construct an example of a matrix A satisfying the following conditions

- 1. The columns of A are 0-1 vectors where the support of no two columns overlap by 50% or more.
- 2. No column's support is totally within the support of another column.
- 3. The minimum 1-norm vector in the column space of A is not a 0-1 vector.

Exercise 10.21 Let M = L + R where L is a low rank matrix corrupted by a sparse noise matrix R. Why can we not recover L from M if R is low rank or if L is sparse?

Exercise 10.22

- 1. Suppose for a univariate convex function f and a finite interval D, $|f''(x)| \leq \delta |f'(x)|$ for every x. Then, what is a good step size to choose for gradient descent? Derive a bound on the number of steps needed to get an approximate minimum of f in terms of as few parameters as possible.
- 2. Generalize the statement and proof to convex functions of d variables.

Exercise 10.23 Prove that the maximum of a convex function over a polytope is attained at one of its vertices.

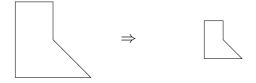
Exercise 10.24 Prove Lemma 10.11.

11 Wavelets

Given a vector space of functions, one would like an orthonormal set of basis functions that span the space. The Fourier transform provides a set of basis functions based on sines and cosines. Often we are dealing with functions that have finite support in which case we would like the basis vectors to have finite support. Also we would like to have an efficient algorithm for computing the coefficients of the expansion of a function in the basis.

11.1 Dilation

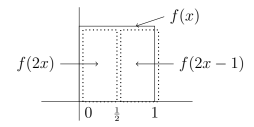
We begin our development of wavelets by first introducing dilation. A *dilation* is a mapping that scales all distances by the same factor.



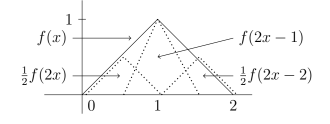
A dilation equation is an equation where a function is defined in terms of a linear combination of scaled, shifted versions of itself. For example,

$$f(x) = \sum_{k=0}^{d-1} c_k f(2x - k).$$

An example is f(x) = f(2x) + f(2x-1) which has a solution f(x) equal one for $0 \le x < 1$ and is zero elsewhere. The equation is illustrated in the figure below. The solid rectangle is f(x) and the dotted rectangles are f(2x) and f(2x-1).



Another example is $f(x) = \frac{1}{2}f(2x) + f(2x-1) + \frac{1}{2}f(2x-2)$. A solution is illustrated in the figure below. The function f(x) is indicated by solid lines. The functions $\frac{1}{2}f(2x)$, f(2x+1), and $\frac{1}{2}f(2x-2)$ are indicated by dotted lines.



Lemma 11.1 If a dilation equation in which all the dilations are a factor of two reduction has a solution, then either the coefficients on the right hand side of the equation sum to two or the integral $\int_{-\infty}^{\infty} f(x) dx$ of the solution is zero.

Proof: Integrate both sides of the dilation equation from $-\infty$ to $+\infty$.

$$\int_{-\infty}^{\infty} f(x)dx = \int_{-\infty}^{\infty} \sum_{k=0}^{d-1} c_k f(2x-k)dx = \sum_{k=0}^{d-1} c_k \int_{-\infty}^{\infty} f(2x-k)dx$$
$$= \sum_{k=0}^{d-1} c_k \int_{-\infty}^{\infty} f(2x)dx = \frac{1}{2} \sum_{k=0}^{d-1} c_k \int_{-\infty}^{\infty} f(x)dx$$

If $\int_{-\infty}^{\infty} f(x) dx \neq 0$, then dividing both sides by $\int_{-\infty}^{\infty} f(x) dx$ gives $\sum_{k=0}^{d-1} c_k = 2$

The above proof interchanged the order of the summation and the integral. This is valid provided the 1-norm of the function is finite. Also note that there are nonzero solutions to dilation equations in which all dilations are a factor of two reduction where the coefficients do not sum to two such as

$$f(x) = f(2x) + f(2x - 1) + f(2x - 2) + f(2x - 3)$$

or

$$f(x) = f(2x) + 2f(2x-1) + 2f(2x-2) + 2f(2x-3) + f(2x-4).$$

In these examples f(x) takes on both positive and negative values and $\int_{-\infty}^{\infty} f(x) dx = 0$.

11.2 The Haar Wavelet

Let $\phi(x)$ be a solution to the dilation equation f(x) = f(2x) + f(2x-1). The function ϕ is called a *scale function* or *scale vector* and is used to generate the two dimensional family of functions, $\phi_{jk} = \phi(2^j x - k)$. Other authors scale $\phi_{jk} = \phi(2^j x - k)$ by $2^{\frac{j}{2}}$ so that the 2-norm, $\int_{-\infty}^{\infty} \phi_{jk}^2(t) dt$, is 1. However, for educational purposes, simplifying the notation for ease of understanding was preferred.

For a given value of j, the shifted versions, $\{\phi_{jk}|k \ge 0\}$, span a space V_j . The spaces V_0, V_1, V_2, \ldots are larger and larger spaces and allow better and better approximations to a function. The fact that $\phi(x)$ is the solution of a dilation equation implies that for for fixed $j \ \phi_{jk}$ is a linear combination of the $\{\phi_{j+1,k}|k \ge 0\}$ and this ensures that $V_j \subseteq V_{j+1}$. It is for this reason that it is desirable in designing a wavelet system for the scale function to satisfy a dilation equation. For a given value of j, the shifted ϕ_{jk} are orthogonal in the sense that $\int_x \phi_{jk}(x)\phi_{jl}(x)dx = 0$ for $k \ne l$.

Note that for each j, the set of functions ϕ_{jk} , k = 0, 1, 2..., form a basis for a vector space V_j and are orthogonal. The set of basis vectors ϕ_{jk} , for all j and k, form an over

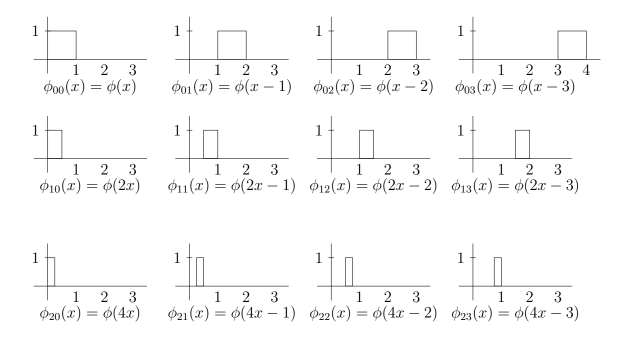
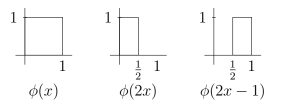


Figure 11.1: Set of scale functions associated with the Haar wavelet.

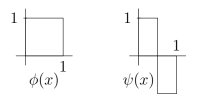
complete basis and for different values of j are not orthogonal. Since ϕ_{jk} , $\phi_{j+1,2k}$, and $\phi_{j+1,2k+1}$ are linearly dependent, for each value of j delete $\phi_{j+1,k}$ for odd values of k to get a linearly independent set of basis vectors. To get an orthogonal set of basis vectors, define

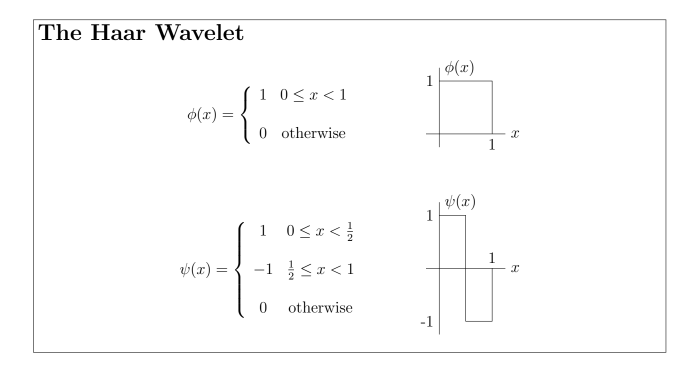
$$\psi_{jk}(x) = \begin{cases} 1 & \frac{2k}{2^{j}} \le x < \frac{2k+1}{2^{j}} \\ -1 & \frac{2k+1}{2^{j}} \le x < \frac{2k+2}{2^{j}} \\ 0 & \text{otherwise} \end{cases}$$

and replace $\phi_{j,2k}$ with $\psi_{j+1,2k}$. Basically, replace the three functions



by the two functions





The basis set becomes

To find a basis for a function that has only finite support, select a scale vector $\phi(x)$ whose scale is that of the support of the function to be represented. Next approximate the function by the set of scale functions $\phi(2^j x - k)$, $k = 0, 1, \ldots$, for some fixed value of j. The value of j is determined by the desired accuracy of the approximation. Basically the x axis has been divided into intervals of size 2^{-j} and in each interval the function is approximated by a fixed value. It is this approximation of the function that is expressed as a linear combination of the basis functions.

Once the value of j has been selected, the function is sampled at 2^j points, one in each interval of width 2^{-j} . Let the sample values be s_0, s_1, \ldots . The approximation to the function is $\sum_{k=0}^{2^j-1} s_k \phi(2^j x - k)$ and is represented by the vector $(s_0, s_1 \ldots, s_{2^j-1})$. The problem now is to represent the approximation to the function using the basis vectors rather than the non orthogonal set of scale functions $\phi_{jk}(x)$. This is illustrated in the following example.

one needs to find the c_i such that

$$\begin{pmatrix} 3\\1\\4\\8\\3\\5\\7\\9 \end{pmatrix} = \begin{pmatrix} 1 & 1 & 1 & 0 & 1 & 0 & 0 & 0\\1 & 1 & 1 & 0 & -1 & 0 & 0 & 0\\1 & 1 & -1 & 0 & 0 & 1 & 0 & 0\\1 & -1 & 0 & 1 & 0 & 0 & -1 & 0\\1 & -1 & 0 & 1 & 0 & 0 & -1 & 0\\1 & -1 & 0 & -1 & 0 & 0 & 0 & 1\\1 & -1 & 0 & -1 & 0 & 0 & 0 & -1 \end{pmatrix} \begin{pmatrix} c_1\\c_2\\c_3\\c_4\\c_5\\c_6\\c_7\\c_8 \end{pmatrix}.$$

The first column represents the scale function $\phi(x)$ and subsequent columns the ψ 's. The tree in Figure 11.2 illustrates an efficient way to find the coefficients representing the vector (3 1 4 8 3 5 7 9) in the basis. Each vertex in the tree contains the average of the quantities of its two children. The root gives the average of the elements in the vector, which is 5 in this example. This average is the coefficient of the basis vector in the first column of the above matrix. The second basis vector converts the average of the eight elements into the average of the first four elements, which is 4, and the last four elements, which is 6, with a coefficient of -1. Working up the tree determines the coefficients for each basis vector.

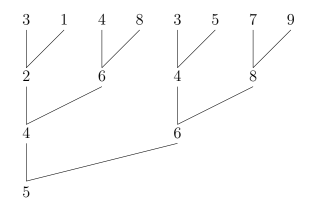


Figure 11.2: Tree of function averages

11.3 Wavelet Systems

So far we have explained wavelets using the simple to understand Haar wavelet. We now consider general wavelet systems. A wavelet system is built from a basic scaling function $\phi(x)$, which comes from a dilation equation. Scaling and shifting of the basic scaling function gives a two dimensional set of scaling functions ϕ_{jk} where

$$\phi_{jk}(x) = \phi(2^j x - k).$$

For a fixed value of j, the ϕ_{jk} span a space V_j . If $\phi(x)$ satisfies a dilation equation

$$\phi(x) = \sum_{k=0}^{d-1} c_k \phi(2x - k),$$

then ϕ_{jk} is a linear combination of the $\phi_{j+1,k}$'s and this implies that $V_0 \subseteq V_1 \subseteq V_2 \subseteq V_3 \cdots$.

11.4 Solving the Dilation Equation

Consider solving a dilation equation

$$\phi(x) = \sum_{k=0}^{d-1} c_k \phi(2x-k)$$

to obtain the scale function for a wavelet system. Perhaps the easiest way is to assume a solution and then calculate the scale function by successive approximation as in the following program for

$$\phi(x) = \frac{1+\sqrt{3}}{4}\phi(2x) + \frac{3+\sqrt{3}}{4}\phi(2x-1) + \frac{3-\sqrt{3}}{4}\phi(2x-2) + \frac{1-\sqrt{3}}{4}\phi(2x-3),$$

a Daubechies scale function. The solution will actually be samples of $\phi(x)$ at some desired resolution.

Set the initial approximation to $\phi(x)$ by generating a vector whose components approximate the samples of $\phi(x)$ at equally spaced values of x.

Calculate the coefficients of the dilation equation.

$$c_1 = \frac{1+\sqrt{3}}{4}$$
 $c_2 = \frac{3+\sqrt{3}}{4}$ $c_3 = \frac{3-\sqrt{3}}{4}$ $c_4 = \frac{1-\sqrt{3}}{4}$

Execute the following loop until the values for $\phi(x)$ converge.

begin

Calculate $\phi(2x)$ by averaging successive values of $\phi(x)$ together. Then fill out the remaining half of the vector representing $\phi(2x)$ with zeros. Calculate $\phi(2x-1)$, $\phi(2x-2)$, and $\phi(2x-3)$ by shifting the contents of $\phi(2x)$ the appropriate distance, discard the zeros that move off the right end and add zeros at the left end.

Calculate the new approximation for $\phi(x)$ using the above values for $\phi(2x-1)$, $\phi(2x-2)$, and $\phi(2x-3)$ in the dilation equation for $\phi(2x)$.

end

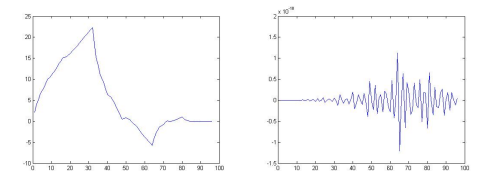


Figure 11.3: Daubechies scale function and associated wavelet

The convergence of the iterative procedure for computing is fast if the eigenvectors of a certain matrix are unity.

Another approach to solving the dilation equation

Consider the dilation equation $\phi(x) = \frac{1}{2}f(2x) + f(2x-1) + \frac{1}{2}f(2x-2)$ and consider continuous solutions with support in $0 \le x < 2$.

$\phi(0) = \frac{1}{2}\phi(0) + \phi(-1) + \phi(-2) = \frac{1}{2}\phi(0) + 0 + 0$	$\phi(0) = 0$
$\phi(2) = \frac{1}{2}\phi(4) + \phi(3) + \phi(2) = \frac{1}{2}\phi(\overline{2}) + 0 + 0$	$\phi(2) = 0$
$\phi(1) = \frac{1}{2}\phi(2) + \phi(1) + \phi(0) = \bar{0} + \phi(1) + 0$	$\phi(1)$ arbitrary

Set $\phi(1) = 1$. Then

$$\phi(\frac{1}{2}) = \frac{1}{2}\phi(1) + \phi(0) + \frac{1}{2}\phi(-1) = \frac{1}{2}$$
$$\phi(\frac{3}{2}) = \frac{1}{2}\phi(3) + \phi(2) + \frac{1}{2}\phi(1) = \frac{1}{2}$$
$$\phi(\frac{1}{4}) = \frac{1}{2}\phi(\frac{1}{2}) + \phi(-\frac{1}{2}) + \frac{1}{2}\phi(-\frac{3}{2}) = \frac{1}{4}$$

Etc.

11.5 Conditions on the Dilation Equation

We would like a basis for a vector space of functions where each basis vector has finite support and the basis vectors are orthogonal. This is achieved by a wavelet system consisting of a shifted version of a scale function that satisfies a dilation equation along with a set of wavelets of various scales and shifts. For the scale function to have a nonzero integral, Lemma 11.1 requires that the coefficients of the dilation equation sum to two. Although the scale function $\phi(x)$ for the Haar system has the property that $\phi(x)$ and $\phi(x-k)$, k > 0, are orthogonal, this is not true for the scale function for the dilation equation $\phi(x) = \frac{1}{2}\phi(2x) + \phi(2x-1) + \frac{1}{2}\phi(2x-2)$. The conditions that integer shifts of the scale function be orthogonal and that the scale function has finite support puts additional conditions on the coefficients of the dilation equation. These conditions are developed in the next two lemmas.

Lemma 11.2 Let

$$\phi(x) = \sum_{k=0}^{d-1} c_k \phi(2x - k).$$

If $\phi(x)$ and $\phi(x-k)$ are orthogonal for $k \neq 0$ and $\phi(x)$ has been normalized so that $\int_{-\infty}^{\infty} \phi(x)\phi(x-k)dx = \delta(k)$, then $\sum_{i=0}^{d-1} c_i c_{i-2k} = 2\delta(k)$.

Proof: Assume $\phi(x)$ has been normalized so that $\int_{-\infty}^{\infty} \phi(x)\phi(x-k)dx = \delta(k)$. Then

$$\int_{x=-\infty}^{\infty} \phi(x)\phi(x-k)dx = \int_{x=-\infty}^{\infty} \sum_{i=0}^{d-1} c_i\phi(2x-i) \sum_{j=0}^{d-1} c_j\phi(2x-2k-j)dx$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} c_ic_j \int_{x=-\infty}^{\infty} \phi(2x-i)\phi(2x-2k-j)dx$$

Since

$$\int_{x=-\infty}^{\infty} \phi(2x-i)\phi(2x-2k-j)dx = \frac{1}{2} \int_{x=-\infty}^{\infty} \phi(y-i)\phi(y-2k-j)dy$$
$$= \frac{1}{2} \int_{x=-\infty}^{\infty} \phi(y)\phi(y+i-2k-j)dy$$
$$= \frac{1}{2}\delta(2k+j-i),$$

 $\int_{x=-\infty}^{\infty} \phi(x)\phi(x-k)dx = \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} c_i c_j \frac{1}{2}\delta(2k+j-i) = \frac{1}{2} \sum_{i=0}^{d-1} c_i c_{i-2k}.$ Since $\phi(x)$ was normalized so that

malized so that

$$\int_{-\infty}^{\infty} \phi(x)\phi(x-k)dx = \delta(k), \text{ it follows that } \sum_{i=0}^{a-1} c_i c_{i-2k} = 2\delta(k).$$

Scale and wavelet coefficients equations

$$\begin{split} \phi(x) &= \sum_{k=0}^{d-1} c_k \phi(2x-k) & \psi(x) = \sum_{k=0}^{d-1} b_k \phi(x-k) \\ \int_{-\infty}^{\infty} \phi(x) \phi(x-k) dx &= \delta(k) & \int_{x=-\infty}^{\infty} \phi(x) \psi(x-k) = 0 \\ \int_{j=0}^{d-1} c_j = 2 & \int_{x=-\infty}^{\infty} \psi(x) dx = 0 \\ \int_{j=0}^{d-1} c_j c_{j-2k} &= 2\delta(k) & \int_{x=-\infty}^{\infty} \psi(x) \psi(x-k) dx = \delta(k) \\ c_k &= 0 \text{ unless } 0 \le k \le d-1 & \sum_{i=0}^{d-1} (-1)^k b_i b_{i-2k} = 2\delta(k) \\ d \text{ even} & \sum_{j=0}^{d-1} c_{2j+1} & \sum_{j=0}^{d-1} c_j b_{j-2k} = 0 \\ \int_{j=0}^{d-1} b_j = 0 \\ b_k &= (-1)^k c_{d-1-k} \end{split}$$

One designs wavelet systems so the above conditions are satisfied.

Lemma 11.2 provides a necessary but not sufficient condition on the coefficients of the dilation equation for shifts of the scale function to be orthogonal. One should note that the conditions of Lemma 11.2 are not true for the triangular or piecewise quadratic solutions to

$$\phi(x) = \frac{1}{2}\phi(2x) + \phi(2x-1) + \frac{1}{2}\phi(2x-2)$$

and

$$\phi(x) = \frac{1}{4}\phi(2x) + \frac{3}{4}\phi(2x-1) + \frac{3}{4}\phi(2x-2) + \frac{1}{4}\phi(2x-3)$$

which overlap and are not orthogonal.

For $\phi(x)$ to have finite support the dilation equation can have only a finite number of terms. This is proved in the following lemma.

Lemma 11.3 If $0 \le x < d$ is the support of $\phi(x)$, and the set of integer shifts, $\{\phi(x - k) | k \ge 0\}$, are linearly independent, then $c_k = 0$ unless $0 \le k \le d - 1$.

Proof: If the support of $\phi(x)$ is $0 \le x < d$, then the support of $\phi(2x)$ is $0 \le x < \frac{d}{2}$. If

$$\phi(x) = \sum_{k=-\infty}^{\infty} c_k \phi(2x - k)$$

the support of both sides of the equation must be the same. Since the $\phi(x-k)$ are linearly independent the limits of the summation are actually k = 0 to d - 1 and

$$\phi(x) = \sum_{k=0}^{d-1} c_k \phi(2x - k).$$

It follows that $c_k = 0$ unless $0 \le k \le d - 1$.

The condition that the integer shifts are linearly independent is essential to the proof and the lemma is not true without this condition.

One should also note that $\sum_{i=0}^{d-1} c_i c_{i-2k} = 0$ for $k \neq 0$ implies that d is even since for d odd and $k = \frac{d-1}{2}$ d-1 d-1

$$\sum_{i=0}^{a-1} c_i c_{i-2k} = \sum_{i=0}^{a-1} c_i c_{i-d+1} = c_{d-1} c_0.$$

For $c_{d-1}c_0$ to be zero either c_{d-1} or c_0 must be zero. Since either $c_0 = 0$ or $c_{d-1} = 0$, there are only d-1 nonzero coefficients. From here on we assume that d is even. If the dilation equation has d terms and the coefficients satisfy the linear equation $\sum_{k=0}^{d-1} c_k = 2$ and the $\frac{d}{2}$ quadratic equations $\sum_{i=0}^{d-1} c_i c_{i-2k} = 2\delta(k)$ for $1 \le k \le \frac{d-1}{2}$, then for d > 2 there are $\frac{d}{2}-1$ coefficients that can be used to design the wavelet system to achieve desired properties.

11.6 Derivation of the Wavelets from the Scaling Function

In a wavelet system one develops a mother wavelet as a linear combination of integer shifts of a scaled version of the scale function $\phi(x)$. Let the mother wavelet $\psi(x)$ be given by $\psi(x) = \sum_{k=0}^{d-1} b_k \phi(2x-k)$. One wants integer shifts of the mother wavelet $\psi(x-k)$ to be orthogonal and also for integer shifts of the mother wavelet to be orthogonal to the scaling function $\phi(x)$. These conditions place restrictions on the coefficients b_k which are the subject matter of the next two lemmas.

Lemma 11.4 (Orthogonality of $\psi(x)$ and $\psi(x-k)$) Let $\psi(x) = \sum_{k=0}^{d-1} b_k \phi(2x-k)$. If $\psi(x)$ and $\psi(x-k)$ are orthogonal for $k \neq 0$ and $\psi(x)$ has been normalized so that $\int_{-\infty}^{\infty} \psi(x)\psi(x-k)dx = \delta(k)$, then

$$\sum_{i=0}^{d-1} (-1)^k b_i b_{i-2k} = 2\delta(k).$$

Proof: Analogous to Lemma 11.2.

Lemma 11.5 (Orthogonality of $\phi(x)$ and $\psi(x-k)$) Let $\phi(x) = \sum_{k=0}^{d-1} c_k \phi(2x-k)$ and $\psi(x) = \sum_{k=0}^{d-1} b_k \phi(2x-k)$. If $\int_{x=-\infty}^{\infty} \phi(x)\phi(x-k)dx = \delta(k)$ and $\int_{x=-\infty}^{\infty} \phi(x)\psi(x-k)dx = 0$ for all k, then $\sum_{i=0}^{d-1} c_i b_{i-2k} = 0$ for all k.

Proof:

$$\int_{x=-\infty}^{\infty} \phi(x)\psi(x-k)dx = \int_{x=-\infty}^{\infty} \sum_{i=0}^{d-1} c_i\phi(2x-i) \sum_{j=1}^{d-1} b_j\phi(2x-2k-j)dx = 0.$$

Interchanging the order of integration and summation

$$\sum_{i=0}^{d-1} \sum_{j=0}^{d-1} c_i b_j \int_{x=-\infty}^{\infty} \phi(2x-i)\phi(2x-2k-j)dx = 0$$

Substituting y = 2x - i yields

$$\frac{1}{2}\sum_{i=0}^{d-1}\sum_{j=0}^{d-1}c_ib_j\int_{y=-\infty}^{\infty}\phi(y)\phi(y-2k-j+i)dy = 0$$

Thus,

$$\sum_{i=0}^{d-1} \sum_{j=0}^{d-1} c_i b_j \delta(2k+j-i) = 0$$

Summing over j gives

$$\sum_{i=0}^{d-1} c_i b_{i-2k} = 0$$

Lemma 11.5 gave a condition on the coefficients in the equations for $\phi(x)$ and $\psi(x)$ if integer shifts of the mother wavelet are to be orthogonal to the scale function. In addition, for integer shifts of the mother wavelet to be orthogonal to the scale function requires that $b_k = (-1)^k c_{d-1-k}$.

Lemma 11.6 Let the scale function $\phi(x)$ equal $\sum_{k=0}^{d-1} c_k \phi(2x-k)$ and let the wavelet function

 $\psi(x) equal \sum_{k=0}^{d-1} b_k \phi(2x-k)$. If the scale functions are orthogonal

$$\int_{-\infty}^{\infty} \phi(x)\phi(x-k)dx = \delta(k)$$

and the wavelet functions are orthogonal with the scale function

$$\int_{x=-\infty}^{\infty} \phi(x)\psi(x-k)dx = 0$$

for all k, then $b_k = (-1)^k c_{d-1-k}$.

Proof: By Lemma 11.5, $\sum_{j=0}^{d-1} c_j b_{j-2k} = 0$ for all k. Separating $\sum_{j=0}^{d-1} c_j b_{j-2k} = 0$ into odd and even indices gives

$$\sum_{j=0}^{\frac{d}{2}-1} c_{2j}b_{2j-2k} + \sum_{j=0}^{\frac{d}{2}-1} c_{2j+1}b_{2j+1-2k} = 0$$
(11.1)

for all k.

$$c_{0}b_{0} + c_{2}b_{2} + c_{4}b_{4} + \dots + c_{1}b_{1} + c_{3}b_{3} + c_{5}b_{5} + \dots = 0 \qquad k = 0$$

$$c_{2}b_{0} + c_{4}b_{2} + \dots + c_{3}b_{1} + c_{5}b_{3} + \dots = 0 \qquad k = 1$$

$$c_{4}b_{0} + \dots + c_{5}b_{1} + \dots = 0 \qquad k = 2$$

By Lemmas 11.2 and 11.4, $\sum_{j=0}^{d-1} c_j c_{j-2k} = 2\delta(k)$ and $\sum_{j=0}^{d-1} b_j b_{j-2k} = 2\delta(k)$ and for all k. Separating odd and even terms,

$$\sum_{j=0}^{\frac{d}{2}-1} c_{2j}c_{2j-2k} + \sum_{j=0}^{\frac{d}{2}-1} c_{2j+1}c_{2j+1-2k} = 2\delta(k)$$
(11.2)

and

$$\sum_{j=0}^{\frac{d}{2}-1} b_{2j}b_{2j-2k} + \sum_{j=0}^{\frac{d}{2}-1} (-1)^j b_{2j+1}b_{2j+1-2k} = 2\delta(k)$$
(11.3)

for all k.

$$c_{0}c_{0} + c_{2}c_{2} + c_{4}c_{4} + \dots + c_{1}c_{1} + c_{3}c_{3} + c_{5}c_{5} + \dots = 2 \qquad k = 0$$

$$c_{2}c_{0} + c_{4}c_{2} + \dots + c_{3}c_{1} + c_{5}c_{3} + \dots = 0 \qquad k = 1$$

$$c_{4}c_{0} + \dots + c_{5}c_{1} + \dots = 0 \qquad k = 2$$

$$b_{0}b_{0} + b_{2}b_{2} + b_{4}b_{4} + \dots + b_{1}b_{1} - b_{3}b_{3} + b_{5}b_{5} - \dots = 2 \qquad k = 0$$

$$b_{2}b_{0} + b_{4}b_{2} + \dots - b_{3}b_{1} + b_{5}b_{3} - \dots = 0 \qquad k = 1$$

$$b_{4}b_{0} + \dots + b_{5}b_{1} - \dots = 0 \qquad k = 2$$

Let $C_e = (c_0, c_2, \ldots, c_{d-2})$, $C_o = (c_1, c_3, \ldots, c_{d-1})$, $B_e = (b_0, b_2, \ldots, b_{d-2})$, and $B_o = (b_1, b_3, \ldots, b_{d-1})$. Equations 12.1, 12.2, and 11.3 can be expressed as convolutions³⁵ of these sequences. Equation 12.1 is $C_e * B_e^R + C_o * B_o^R = 0$, 12.2 is $C_e * C_e^R + C_o * C_o^R = \delta(k)$, and 11.3 is $B_e * B_e^R + B_o * B_o^R = \delta(k)$, where the superscript R stands for reversal of the sequence. These equations can be written in matrix format as

$$\begin{pmatrix} C_e & C_o \\ B_e & B_o \end{pmatrix} * \begin{pmatrix} C_e^R & B_e^R \\ C_o^R & B_o^R \end{pmatrix} = \begin{pmatrix} 2\delta & 0 \\ 0 & 2\delta \end{pmatrix}$$

Taking the Fourier or z-transform yields

$$\left(\begin{array}{cc} F(C_e) & F(C_o) \\ F(B_e) & F(B_o) \end{array}\right) \left(\begin{array}{cc} F(C_e^R) & F(B_e^R) \\ F(C_o^R) & F(B_o^R) \end{array}\right) = \left(\begin{array}{cc} 2 & 0 \\ 0 & 2 \end{array}\right).$$

where F denotes the transform. Taking the determinant yields

$$\left(F(C_e)F(B_o) - F(B_e)F(C_o)\right)\left(F(C_e)F(B_o) - F(C_o)F(B_e)\right) = 4$$

Thus $F(C_e)F(B_o) - F(C_o)F(B_e) = 2$ and the inverse transform yields

$$C_e * B_o - C_o * B_e = 2\delta(k).$$

Convolution by C_e^R yields

$$C_e^R * C_e * B_o - C_e^R * B_e * C_o = C_e^R * 2\delta(k)$$

Now
$$\sum_{j=0}^{d-1} c_j b_{j-2k} = 0$$
 so $-C_e^R * B_e = C_o^R * B_o$. Thus
 $C_e^R * C_e * B_o + C_o^R * B_o * C_o = 2C_e^R * \delta(k)$
 $(C_e^R * C_e + C_o^R * C_o) * B_o = 2C_e^R * \delta(k)$
 $2\delta(k) * B_o = 2C_e^R * \delta(k)$
 $C_e = B_o^R$

Thus, $c_i = 2b_{d-1-i}$ for even *i*. By a similar argument, convolution by C_0^R yields

$$C_0^R * C_e * B_0 - C_0^R * C_0 * B_e = 2C_0^R \delta(k)$$

Since $C_{)}^{R} * B_{0} = -C_{0}^{R} * B_{e}$

$$-C_{e}^{R} * C_{e}^{R} * B_{e} - C_{0}^{R} * C_{0} * B_{e} = 2C_{0}^{R}\delta(k)$$

-(C_{e} * C_{e}^{R} + C_{0}^{R} * C_{0}) * B_{e} = 2C_{0}^{R}\delta(k)
-2\delta(k)B_{e} = 2C_{0}^{R}\delta(k)
-B_{e} = C_{0}^{R}

Thus, $c_i = -2b_{d-1-i}$ for all odd *i* and hence $c_i = (-1)^i 2b_{d-1-i}$ for all *i*.

 $[\]overline{ (a_0, a_1, \dots, a_{d-1}) }$ and $(b_0, b_1, \dots, b_{d-1})$ denoted $(a_0, a_1, \dots, a_{d-1}) * (b_0, b_1, \dots, b_{d-1})$ is the sequence $(a_0b_{d-1}, a_0b_{d-2} + a_1b_{d-1}, a_0b_{d-3} + a_1b_{d-2} + a_3b_{d-1}\dots, a_{d-1}b_0).$

11.7 Sufficient Conditions for the Wavelets to be Orthogonal

Section 11.6 gave necessary conditions on the b_k and c_k in the definitions of the scale function and wavelets for certain orthogonality properties. In this section we show that these conditions are also sufficient for certain orthogonality conditions. One would like a wavelet system to satisfy certain conditions.

- 1. Wavelets, $\psi_j(2^j x k)$, at all scales and shifts to be orthogonal to the scale function $\phi(x)$.
- 2. All wavelets to be orthogonal. That is

$$\int_{-\infty}^{\infty} \psi_j (2^j x - k) \psi_l (2^l x - m) dx = \delta(j - l) \delta(k - m)$$

3. $\phi(x)$ and ψ_{jk} , $j \leq l$ and all k, to span V_l , the space spanned by $\phi(2^l x - k)$ for all k. These items are proved in the following lemmas. The first lemma gives sufficient conditions on the wavelet coefficients b_k in the definition

$$\psi(x) = \sum_{k} b_k \psi(2x - k)$$

for the mother wavelet so that the wavelets will be orthogonal to the scale function. That is, if the wavelet coefficients equal the scale coefficients in reverse order with alternating negative signs, then the wavelets will be orthogonal to the scale function.

Lemma 11.7 If $b_k = (-1)^k c_{d-1-k}$, then $\int_{-\infty}^{\infty} \phi(x) \psi(2^j x - l) dx = 0$ for all j and l.

Proof: Assume that $b_k = (-1)^k c_{d-1-k}$. We first show that $\phi(x)$ and $\psi(x-k)$ are orthogonal for all values of k. Then we modify the proof to show that $\phi(x)$ and $\psi(2^j x - k)$ are orthogonal for all j and k.

Assume $b_k = (-1)^k c_{d-1-k}$. Then

$$\int_{-\infty}^{\infty} \phi(x)\psi(x-k) = \int_{-\infty}^{\infty} \sum_{i=0}^{d-1} c_i \phi(2x-i) \sum_{j=0}^{d-1} b_j \phi(2x-2k-j) dx$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} c_i (-1)^j c_{d-1-j} \int_{-\infty}^{\infty} \phi(2x-i) \phi(2x-2k-j) dx$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} (-1)^j c_i c_{d-1-j} \delta(i-2k-j)$$
$$= \sum_{j=0}^{d-1} (-1)^j c_{2k+j} c_{d-1-j}$$
$$= c_{2k} c_{d-1} - c_{2k+1} c_{d-2} + \dots + c_{d-2} c_{2k-1} - c_{d-1} c_{2k}$$
$$= 0$$

The last step requires that d be even which we have assumed for all scale functions.

For the case where the wavelet is $\psi(2^j - l)$, first express $\phi(x)$ as a linear combination of $\phi(2^{j-1}x - n)$. Now for each these terms

$$\int_{-\infty}^{\infty} \phi(2^{j-1}x - m)\psi(2^jx - k)dx = 0$$

To see this, substitute $y = 2^{j-1}x$. Then

$$\int_{-\infty}^{\infty} \phi(2^{j}x - m)\psi(2^{j}x - k)dx = \frac{1}{2^{j-1}}\int_{-\infty}^{\infty} \phi(y - m)\psi(2y - k)dy$$

which by the previous argument is zero.

The next lemma gives conditions on the coefficients b_k that are sufficient for the wavelets to be orthogonal.

Lemma 11.8 If $b_k = (-1)^k c_{d-1-k}$, then

$$\int_{-\infty}^{\infty} \frac{1}{2^j} \psi_j (2^j x - k) \frac{1}{2^k} \psi_l (2^l x - m) dx = \delta(j - l) \delta(k - m).$$

Proof: The first level wavelets are orthogonal.

$$\int_{-\infty}^{\infty} \psi(x)\psi(x-k)dx = \int_{-\infty}^{\infty} \sum_{i=0}^{d-1} b_i \phi(2x-i) \sum_{j=0}^{d-1} b_j \phi(2x-2k-j)dx$$
$$= \sum_{i=0}^{d-1} b_i \sum_{j=0}^{d-1} b_j \int_{-\infty}^{\infty} \phi(2x-i)\phi(2x-2k-j)dx$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} b_i b_j \delta(i-2k-j)$$
$$= \sum_{i=0}^{d-1} bib_{i-2k}$$
$$= \sum_{i=0}^{d-1} (-1)^i c_{d-1-i} (-1)^{i-2k} c_{d-1-i+2k}$$
$$= \sum_{i=0}^{d-1} (-1)^{2i-2k} c_{d-1-i} c_{d-1-i+2k}$$

Substituting j for d - 1 - i yields

$$\sum_{j=0}^{d-1} c_j c_{j+2k} = 2\delta(k)$$

Example of orthogonality when wavelets are of different scale.

$$\int_{-\infty}^{\infty} \psi(2x)\psi(x-k)dx = \int_{-\infty}^{\infty} \sum_{i=0}^{d-1} b_i \phi(4x-i) \sum_{j=0}^{d-1} b_j \phi(2x-2k-j)dx$$
$$= \sum_{i=0}^{d-1} \sum_{i=0}^{d-1} b_i b_j \int_{-\infty}^{\infty} \phi(4x-i)\phi(2x-2k-j)dx$$

Since $\phi(2x - 2k - j) = \sum_{l=0}^{d-1} c_l \phi(4x - 4k - 2j - l)$

$$\int_{-\infty}^{\infty} \psi(2x)\psi(x-k)dx = \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} \sum_{l=0}^{d-1} b_i b_j c_l \int_{-\infty}^{\infty} \psi(4x-i)\phi(4x-4k-2j-l)dx$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} \sum_{l=0}^{d-1} b_i b_j c_l \delta(i-4k-2j-l)$$
$$= \sum_{i=0}^{d-1} \sum_{j=0}^{d-1} b_i b_j c_{i-4k-2j}$$

Since $\sum_{j=0}^{d-1} c_j b_{j-2k} = 0$, $\sum_{i=0}^{d-1} b_i c_{i-4k-2j} = \delta(j-2k)$ Thus

$$\int_{-\infty}^{\infty} \psi(2x)\psi(x-k)dx = \sum_{j=0}^{d-1} b_j \delta(j-2k) = 0.$$

Orthogonality of scale function with wavelet of different scale.

$$\int_{-\infty}^{\infty} \phi(x)\psi(2x-k)dx = \int_{-\infty}^{\infty} \sum_{j=0}^{d-1} c_j\phi(2x-j)\psi(2x-k)dx$$
$$= \sum_{j=0}^{d-1} c_j \int_{-\infty}^{\infty} \phi(2x-j)\psi(2x-k)dx$$
$$= \frac{1}{2} \sum_{j=0}^{d-1} c_j \int_{-\infty}^{\infty} \phi(y-j)\psi(y-k)dy$$
$$= 0$$

If ψ was of scale 2^j , ϕ would be expanded as a linear combination of ϕ of scale 2^j all of which would be orthogonal to ψ .

11.8 Expressing a Function in Terms of Wavelets

Given a wavelet system with scale function ϕ and mother wavelet ψ we wish to express a function f(x) in terms of an orthonormal basis of the wavelet system. First we will express f(x) in terms of scale functions $\phi_{jk}(x) = \phi(2^j x - k)$. To do this we will build a tree similar to that in Figure 11.2 for the Haar system only computing the coefficients will be much more complex. Recall that the coefficients at a level in the tree are the coefficients to represent f(x) using scale functions with the precision of the level.

Let $f(x) = \sum_{k=0}^{\infty} a_{jk} \phi_j(x-k)$ where the a_{jk} are the coefficients in the expansion of f(x) using level j scale functions. Since the $\phi_j(x-k)$ are orthogonal

$$a_{jk} = \int_{x=-\infty}^{\infty} f(x)\phi_j(x-k)dx.$$

Expanding ϕ_j in terms of ϕ_{j+1} yields

$$a_{jk} = \int_{x=-\infty}^{\infty} f(x) \sum_{m=0}^{d-1} c_m \phi_{j+1} (2x - 2k - m) dx$$
$$= \sum_{m=0}^{d-1} c_m \int_{x=-\infty}^{\infty} f(x) \phi_{j+1} (2x - 2k - m) dx$$
$$= \sum_{m=0}^{d-1} c_m a_{j+1,2k+m}$$

Let n = 2k + m. Now m = n - 2k. Then

$$a_{jk} = \sum_{n=2k}^{d-1} c_{n-2k} a_{j+1,n} \tag{11.4}$$

In construction the tree similar to that in Figure 11.2, the values at the leaves are the values of the function sampled in the intervals of size 2^{-j} . Equation 11.4 is used to compute values as one moves up the tree. The coefficients in the tree could be used if we wanted to represent f(x) using scale functions. However, we want to represent f(x) using one scale function whose scale is the support of f(x) along with wavelets which gives us an orthogonal set of basis functions. To do this we need to calculate the coefficients for the wavelets.. The value at the root of the tree is the coefficient for the scale function. We then move down the tree calculating the coefficients for the wavelets.

Finish by calculating wavelet coefficients maybe add material on jpeg

Example: Add example using D_4 . Maybe example using sinc

11.9 Designing a Wavelet System

In designing a wavelet system there are a number of parameters in the dilation equation. If one uses d terms in the dilation equation, one degree of freedom can be used to satisfy

$$\sum_{i=0}^{d-1} c_i = 2$$

which insures the existence of a solution with a nonzero mean. Another $\frac{d}{2}$ degrees of freedom are used to satisfy

$$\sum_{i=0}^{d-1} c_i c_{i-2k} = \delta(k)$$

which insures the orthogonal properties. The remaining $\frac{d}{2} - 1$ degrees of freedom can be used to obtain some desirable properties such as smoothness. Smoothness appears to be related to vanishing moments of the scaling function. Material on the design of systems is beyond the scope of this book and can be found in the literature.

Exercises

Exercise 11.1 What is the solution to the dilation equation f(x) = f(2x) + f(2x - k) for k an integer?

Exercise 11.2 Are there solutions to f(x) = f(2x) + f(2x - 1) other than a constant multiple of

$$f(x) = \begin{cases} 1 & 0 \le x < 1 \\ 0 & otherwise \end{cases}$$
?

Exercise 11.3 Is there a solution to $f(x) = \frac{1}{2}f(2x) + f(2x-1) + \frac{1}{2}f(2x-2)$ with f(0) = f(1) = 1 and f(2) = 0?

Exercise 11.4 What is the solution to the dilation equation

$$f(x) = f(2x) + f(2x - 1) + f(2x - 2) + f(2x - 3).$$

Exercise 11.5 Consider the dilation equation

$$f(x) = f(2x) + 2f(2x - 1) + 2f(2x - 2) + 2f(2x - 3) + f(2x - 4)$$

- 1. What is the solution to the dilation equation?
- 2. What is the value of $\int_{-\infty}^{\infty} f(x) dx$?

Exercise 11.6 What are the solutions to the following families of dilation equations.

1.

$$\begin{aligned} f(x) &= f(2x) + f(2x-1) \\ f(x) &= \frac{1}{2}f(2x) + \frac{1}{2}f(2x-1) + \frac{1}{2}f(2x-2) + \frac{1}{2}f(2x-3) \\ f(x) &= \frac{1}{4}f(2x) + \frac{1}{4}f(2x-1) + \frac{1}{4}f(2x-2) + \frac{1}{4}f(2x-3) + \frac{1}{4}f(2x-4) + \frac{1}{4}f(2x-5) \\ &+ \frac{1}{4}f(2x-6) + \frac{1}{4}f(2x-7) \\ f(x) &= \frac{1}{k}f(2x) + \frac{1}{k}f(2x) + \dots + \frac{1}{k}f(2x) \end{aligned}$$

2.

$$\begin{split} f(x) &= \frac{1}{3}f(2x) + \frac{2}{3}f(2x-1) + \frac{2}{3}f(2x-2) + \frac{1}{3}f(2x-3) \\ f(x) &= \frac{1}{4}f(2x) + \frac{3}{4}f(2x-1) + \frac{3}{4}f(2x-2) + \frac{1}{4}f(2x-3) \\ f(x) &= \frac{1}{5}f(2x) + \frac{4}{5}f(2x-1) + \frac{4}{5}f(2x-2) + \frac{1}{5}f(2x-3) \\ f(x) &= \frac{1}{k}f(2x) + \frac{k-1}{k}f(2x-1) + \frac{k-1}{k}f(2x-2) + \frac{1}{k}f(2x-3) \end{split}$$

3.

$$\begin{aligned} f(x) &= \frac{1}{2}f(2x) + \frac{1}{2}f(2x-1) + \frac{1}{2}f(2x-2) + \frac{1}{2}f(2x-3) \\ f(x) &= \frac{3}{2}f(2x) - \frac{1}{2}f(2x-1) + \frac{3}{2}f(2x-2) - \frac{1}{2}f(2x-3) \\ f(x) &= \frac{5}{2}f(2x) - \frac{3}{2}f(2x-1) + \frac{5}{2}f(2x-2) - \frac{3}{2}f(2x-3) \\ f(x) &= \frac{1+2k}{2}f(2x) - \frac{2k-1}{2}f(2x-1) + \frac{1+2k}{2}f(2x-2) - \frac{2k-1}{2}f(2x-3) \end{aligned}$$

4.

$$\begin{split} f(x) &= \frac{1}{3}f(2x) + \frac{2}{3}f(2x-1) + \frac{2}{3}f(2x-2) + \frac{1}{3}f(2x-3) \\ f(x) &= \frac{4}{3}f(2x) - \frac{1}{3}f(2x-1) + \frac{5}{3}f(2x-2) - \frac{2}{3}f(2x-3) \\ f(x) &= \frac{7}{3}f(2x) - \frac{4}{3}f(2x-1) + \frac{8}{3}f(2x-2) - \frac{5}{3}f(2x-3) \\ f(x) &= \frac{1+3k}{3}f(2x) - \frac{2-3k}{3}f(2x-1) + \frac{2+3k}{3}f(2x-2) - \frac{1-3k}{3}f(2x-3) \end{split}$$

Exercise 11.7

Solution:

Exercise 11.8

- 1. What is the solution to the dilation equation $f(x) = \frac{1}{2}f(2x) + \frac{3}{2}f(2x-1)$? Hint: Write a program to see what the solution looks like.
- 2. How does the solution change when the equation is changed to $f(x) = \frac{1}{3}f(2x) + \frac{5}{3}f(2x-1)?$
- 3. How does the solution change if the coefficients no longer sum to two as in f(x) = f(2x) + 3f(2x-1)?

Exercise 11.9 If f(x) is frequency limited by 2π , prove that

$$f(x) = \sum_{k=0}^{\infty} f(k) \frac{\sin(\pi(x-k))}{\pi(x-k)}.$$

Hint: Use the Nyquist sampling theorem which states that a function frequency limited by 2π is completely determined by samples spaced one unit apart. Note that this result means that

$$f(k) = \int_{-\infty}^{\infty} f(x) \frac{\sin(\pi(x-k))}{\pi(x-k)} dx$$

Exercise 11.10 Compute an approximation to the scaling function that comes from the dilation equation

$$\phi(x) = \frac{1+\sqrt{3}}{4}\phi(2x) + \frac{3+\sqrt{3}}{4}\phi(2x-1) + \frac{3-\sqrt{3}}{4}\phi(2x-2) + \frac{1\sqrt{3}}{4}\phi(2x-3).$$

Exercise 11.11 Consider f(x) to consist of the semi circle $(x - \frac{1}{2})^2 + y^2 = \frac{1}{4}$ and $y \ge 0$ for $0 \le x \le 1$ and 0 otherwise.

1. Using precision j = 4 find the coefficients for the scale functions and the wavelets for D_4 defined by the dilation equation

$$\phi(x) = \frac{1+\sqrt{3}}{4}\phi(2x) + \frac{3+\sqrt{3}}{4}\phi(2x-1) + \frac{3-\sqrt{3}}{4}\phi(2x-2) + \frac{1\sqrt{3}}{4}\phi(2x-3)$$

2. Graph the approximation to the semi circle for precision j = 4.

Exercise 11.12 What is the set of all solutions to the dilation equation

$$\phi(x) = \frac{1+\sqrt{3}}{4}\phi(2x) + \frac{3+\sqrt{3}}{4}\phi(2x-1) + \frac{3-\sqrt{3}}{4}\phi(2x-2) + \frac{1\sqrt{3}}{4}\phi(2x-3)$$

Exercise 11.13 Prove that if scale functions defined by a dilation equation are orthogonal, then the sum of the even coefficients must equal the sum of the odd coefficients in the dilation equation. That is, $\sum_{k} c_{2k} = \sum_{k} c_{2k+1}$.

```
function = wavelets
```

```
acc=32; %accuracy of computation
phit=[1:acc zeros(1,3*acc)];
```

c1=(1+3^0.5)/4; c2=(3+3^0.5)/4; c3=(3-3^0.5)/4; c4=(1-3^0.5)/4;

```
for i=1:10
   temp=(phit(1:2:4*acc)+phit(2:2:4*acc))/2;
   phi2t=[temp zeros(1,3*acc)];
   phi2tshift1=[ zeros(1,acc) temp zeros(1,2*acc)];
   phi2tshift2=[ zeros(1,2*acc) temp zeros(1,acc)];
   phi2tshift3=[ zeros(1,3*acc) temp ];
   phit=c1*phi2t+c2*phi2tshift1+c3*phi2tshift2+c4*phi2tshift3;
   plot(phit)
   figure(gcf)
   pause
end plot(phit) figure(gcf) end
```

12 Appendix

12.1 Asymptotic Notation

We introduce the big O notation here. The motivating example is the analysis of the running time of an algorithm. The running time may be a complicated function of the input length n such as $5n^3 + 25n^2 \ln n - 6n + 22$. Asymptotic analysis is concerned with the behavior as $n \to \infty$ where the higher order term $5n^3$ dominates. Further, the coefficient 5 of $5n^3$ is not of interest since its value varies depending on the machine model. So we say that the function is $O(n^3)$. The big O notation applies to functions on the positive integers taking on positive real values.

Definition 12.1 For functions f and g from the natural numbers to the positive reals, f(n) is O(g(n)) if there exists a constant c > 0 such that for all n, $f(n) \le cg(n)$.

Thus, $f(n) = 5n^3 + 25n^2 \ln n - 6n + 22$ is $O(n^3)$. The upper bound need not be tight. Not only is f(n), $O(n^3)$, it is also $O(n^4)$. Note g(n) must be strictly greater than 0 for all n.

To say that the function f(n) grows at least as fast as g(n), one uses a notation called omega of n. For positive real valued f and g, f(n) is $\Omega(g(n))$ if there exists a constant c > 0 such that for all n, $f(n) \ge cg(n)$. If f(n) is both O(g(n)) and $\Omega(g(n))$, then f(n) is $\Theta(g(n))$. Theta of n is used when the two functions have the same asymptotic growth rate.

Many times one wishes to bound the low order terms. To do this, a notation called little o of n is used. We say f(n) is o(g(n)) if $\lim_{n\to\infty} \frac{f(n)}{g(n)} = 0$. Note that f(n) being O(g(n)) means that asymptotically f(n) does not grow faster than g(n), whereas f(n)being o(g(n)) means that asymptotically f(n)/g(n) goes to zero. If $f(n) = 2n + \sqrt{n}$, then

 $\begin{array}{ll} \textbf{asymptotic upper bound} \\ f(n) \text{ is } O(g(n)) \text{ if for all } n, f(n) \leq cg(n) \text{ for some constant } c > 0. \\ \end{array} \leq \\ \textbf{asymptotic lower bound} \\ f(n) \text{ is } \Omega(g(n)) \text{ if for all } n, f(n) \geq cg(n) \text{ for some constant } c > 0. \\ \end{array} \geq \\ \textbf{asymptotic equality} \\ f(n) \text{ is } \Theta(g(n)) \text{ if it is both } O(g(n)) \text{ and } \Omega(g(n)). \\ = \\ f(n) \text{ is } o(g(n)) \text{ if } \lim_{n \to \infty} \frac{f(n)}{g(n)} = 0. \\ < \\ f(n) \sim g(n) \text{ if } \lim_{n \to \infty} \frac{f(n)}{g(n)} = 1. \\ = \\ f(n) \text{ is } \omega(g(n)) \text{ if } \lim_{n \to \infty} \frac{f(n)}{g(n)} = \infty. \\ \end{array} >$

f(n) is O(n) but in bounding the lower order term, we write f(n) = 2n + o(n). Finally, we write $f(n) \sim g(n)$ if $\lim_{n \to \infty} \frac{f(n)}{g(n)} = 1$ and say f(n) is $\omega(g(n))$ if $\lim_{n \to \infty} \frac{f(n)}{g(n)} = \infty$. The difference between f(n) being $\Theta(g(n))$ and $f(n) \sim g(n)$ is that in the first case f(n) and g(n) may differ by a multiplicative constant factor.

Useful relations 12.2

Summations

$$\begin{split} \sum_{i=0}^{n} a^{i} &= 1 + a + a^{2} + \dots = \frac{1 - a^{n+1}}{1 - a}, \quad a \neq 1 \\ \sum_{i=0}^{\infty} a^{i} &= 1 + a + a^{2} + \dots = \frac{1}{1 - a}, \quad |a| < 1 \\ \sum_{i=0}^{\infty} ia^{i} &= a + 2a^{2} + 3a^{3} \dots = \frac{a}{(1 - a)^{2}}, \quad |a| < 1 \\ \sum_{i=0}^{\infty} i^{2}a^{i} &= a + 4a^{2} + 9a^{3} \dots = \frac{a(1 + a)}{(1 - a)^{3}}, \quad |a| < 1 \\ \sum_{i=1}^{n} i &= \frac{n(n + 1)}{2} \\ \sum_{i=1}^{n} i^{2} &= \frac{n(n + 1)(2n + 1)}{6} \\ \sum_{i=1}^{\infty} \frac{1}{i^{2}} &= \frac{\pi^{2}}{6} \end{split}$$

We prove one equality.

$$\sum_{i=0}^{\infty} ia^i = a + 2a^2 + 3a^3 \dots = \frac{a}{(1-a)^2}, \text{ provided } |a| < 1$$

Write $S = \sum_{i=0}^{\infty} ia^i$.
$$aS = \sum_{i=0}^{\infty} ia^{i+1} = \sum_{i=1}^{\infty} (i-1)a^i.$$
Thus

Thus,

$$S - aS = \sum_{i=1}^{\infty} ia^{i} - \sum_{i=1}^{\infty} (i-1)a^{i} = \sum_{i=1}^{\infty} a^{i} = \frac{a}{1-a},$$

from which the equality follows. The sum $\sum i^2 a^i$ can also be done by an extension of this method (left to the reader). Using generating functions, we will see another proof of both these equalities by derivatives.

$$\sum_{i=1}^{\infty} \frac{1}{i} = 1 + \frac{1}{2} + \left(\frac{1}{3} + \frac{1}{4}\right) + \left(\frac{1}{5} + \frac{1}{6} + \frac{1}{7} + \frac{1}{8}\right) + \dots \ge 1 + \frac{1}{2} + \frac{1}{2} + \dots$$
 and thus diverges.

The summation $\sum_{i=1}^{n} \frac{1}{i}$ grows as $\ln n$ since $\sum_{i=1}^{n} \frac{1}{i} \approx \int_{x=1}^{n} \frac{1}{x} dx$. In fact, $\lim_{i \to \infty} \left(\sum_{i=1}^{n} \frac{1}{i} - \ln(n) \right) = \gamma$ where $\gamma \approx 0.5772$ is Euler's constant. Thus, $\sum_{i=1}^{n} \frac{1}{i} \approx \ln(n) + \gamma$ for large n.

Truncated Taylor series

If all the derivatives of a function f(x) exist, then we can write

$$f(x) = f(0) + f'(0)x + f''(0)\frac{x^2}{2} + \cdots$$

The series can be truncated. In fact, there exists some y between 0 and x such that

$$f(x) = f(0) + f'(y)x.$$

Also, there exists some z between 0 and x such that

$$f(x) = f(0) + f'(0)x + f''(z)\frac{x^2}{2}$$

and so on for higher derivatives. This can be used to derive inequalities. For example, if $f(x) = \ln(1+x)$, then its derivatives are

$$f'(x) = \frac{1}{1+x}; \ f''(x) = -\frac{1}{(1+x)^2}; \ f'''(x) = \frac{2}{(1+x)^3};$$

For any z, f''(z) < 0 and thus for any x, $f(x) \le f(0) + f'(0)x$ hence, $\ln(1+x) \le x$, which also follows from the inequality $1 + x \le e^x$. Also using

$$f(x) = f(0) + f'(0)x + f''(0)\frac{x^2}{2} + f'''(z)\frac{x^3}{3!}$$

for z > -1, f'''(z) > 0, and so for x > -1,

$$\ln(1+x) > x - \frac{x^2}{2}$$

Exponentials and logs

$$a^{\log b} = b^{\log a}$$

 $e^x = 1 + x + \frac{x^2}{2!} + \frac{x^3}{3!} + \dots \qquad e = 2.7182 \qquad \frac{1}{e} = 0.3679$

Setting x = 1 in the equation $e^x = 1 + x + \frac{x^2}{2!} + \frac{x^3}{3!} + \cdots$ yields $e = \sum_{i=0}^{\infty} \frac{1}{i!}$.

$$\lim_{n \to \infty} \left(1 + \frac{a}{n} \right)^n = e^a$$
$$\ln(1+x) = x - \frac{1}{2}x^2 + \frac{1}{3}x^3 - \frac{1}{4}x^4 \cdots \quad |x| < 1$$

The above expression with -x substituted for x gives rise to the approximations

$$\ln(1-x) < -x$$

which also follows from $1 - x \le e^{-x}$, since $\ln(1 - x)$ is a monotone function for $x \in (0, 1)$.

For 0 < x < 0.69, $\ln(1-x) > -x - x^2$.

Trigonometric identities

$$e^{ix} = \cos(x) + i\sin(x)$$

$$\cos(x) = \frac{1}{2} (e^{ix} + e^{-ix})$$

$$\sin(x) = \frac{1}{2i} (e^{ix} - e^{-ix})$$

$$\sin(x \pm y) = \sin(x)\cos(y) \pm \cos(x)\sin(y)$$

$$\cos(x \pm y) = \cos(x)\cos(y) \mp \sin(x)\sin(y)$$

$$\cos(2\theta) = \cos^2 \theta - \sin^2 \theta = 1 - 2\sin^2 \theta$$

$$\sin(2\theta) = 2\sin\theta\cos\theta$$

$$\sin^2 \frac{\theta}{2} = \frac{1}{2} (1 - \cos\theta)$$

$$\cos^2 \frac{\theta}{2} = \frac{1}{2} (1 + \cos\theta)$$

Gaussian and related integrals

$$\int xe^{ax^2} dx = \frac{1}{2a}e^{ax^2}$$

$$\int \frac{1}{a^2 + x^2} dx = \frac{1}{a} \tan^{-1} \frac{x}{a} \text{ thus } \int_{-\infty}^{\infty} \frac{1}{a^2 + x^2} dx = \frac{\pi}{a}$$

$$\int_{-\infty}^{\infty} e^{-\frac{a^2x^2}{2}} dx = \frac{\sqrt{2\pi}}{a} \text{ thus } \frac{a}{\sqrt{2\pi}} \int_{-\infty}^{\infty} e^{-\frac{a^2x^2}{2}} dx = 1$$

$$\int_{0}^{\infty} x^2 e^{-ax^2} dx = \frac{1}{4a} \sqrt{\frac{\pi}{a}}$$

$$\int_{0}^{\infty} x^{2n} e^{-\frac{x^2}{a^2}} dx = \sqrt{\pi} \frac{1 \cdot 3 \cdot 5 \cdots (2n-1)}{2^{n+1}} a^{2n-1} = \sqrt{\pi} \frac{(2n)!}{n!} \left(\frac{a}{2}\right)^{2n+1}$$

$$\int_{0}^{\infty} x^{2n+1} e^{-\frac{x^2}{a^2}} dx = \frac{n!}{2} a^{2n+2}$$

$$\int_{0}^{\infty} e^{-x^2} dx = \sqrt{\pi}$$

To verify $\int_{-\infty}^{\infty} e^{-x^2} dx = \sqrt{\pi}$, consider $\left(\int_{-\infty}^{\infty} e^{-x^2} dx\right)^2 = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} e^{-(x^2+y^2)} dx dy$. Let $x = r \cos \theta$ and $y = r \sin \theta$. The Jacobian of this transformation of variables is

$$J(r,\theta) = \begin{vmatrix} \frac{\partial x}{\partial r} & \frac{\partial x}{\partial \theta} \\ \frac{\partial y}{\partial r} & \frac{\partial y}{\partial \theta} \end{vmatrix} = \begin{vmatrix} \cos \theta & -r\sin \theta \\ \sin \theta & r\cos \theta \end{vmatrix} = r$$

Thus,

$$\left(\int_{-\infty}^{\infty} e^{-x^2} dx\right)^2 = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} e^{-\left(x^2 + y^2\right)} dx dy = \int_{0}^{\infty} \int_{0}^{2\pi} e^{-r^2} J\left(r,\theta\right) dr d\theta$$
$$= \int_{0}^{\infty} e^{-r^2} r dr \int_{0}^{2\pi} d\theta$$
$$= -2\pi \left[\frac{e^{-r^2}}{2}\right]_{0}^{\infty} = \pi$$

Thus, $\int_{-\infty}^{\infty} e^{-x^2} dx = \sqrt{\pi}.$

Miscellaneous integrals

$$\int_{x=0}^{1} x^{\alpha-1} (1-x)^{\beta-1} dx = \frac{\Gamma(\alpha)\Gamma(\beta)}{\Gamma(\alpha+\beta)}$$

For definition of the gamma function see Section 12.3 Binomial coefficients

The binomial coefficient $\binom{n}{k} = \frac{n!}{(n-k)!k!}$ is the number of ways of choosing k items from n. The number of ways of choosing d+1 items from n+1 items equals the number of ways of choosing the d+1 items from the first n items plus the number of ways of choosing d of the items from the first n items with the other item being the last of the n+1 items.

$$\binom{n}{d} + \binom{n}{d+1} = \binom{n+1}{d+1}.$$

The observation that the number of ways of choosing k items from 2n equals the number of ways of choosing i items from the first n and choosing k - i items from the second n summed over all $i, 0 \le i \le k$ yields the identity

$$\sum_{i=0}^{k} \binom{n}{i} \binom{n}{k-i} = \binom{2n}{k}.$$

Setting k = n in the above formula and observing that $\binom{n}{i} = \binom{n}{n-i}$ yields

$$\sum_{i=0}^{n} \binom{n}{i}^2 = \binom{2n}{n}.$$

More generally $\sum_{i=0}^{k} {n \choose i} {m \choose k-i} = {n+m \choose k}$ by a similar derivation.

12.3 Useful Inequalities

 $1 + x \le e^x$ for all real x.

One often establishes an inequality such as $1 + x \leq e^x$ by showing that the difference of the two sides, namely $e^x - (1 + x)$, is always positive. This can be done by taking derivatives. The first and second derivatives are $e^x - 1$ and e^x . Since e^x is always positive, $e^x - 1$ is monotonic and $e^x - (1 + x)$ is convex. Since $e^x - 1$ is monotonic, it can be zero only once and is zero at x = 0. Thus, $e^x - (1 + x)$ takes on its minimum at x = 0 where it is zero establishing the inequality.

$$(1-x)^n \ge 1 - nx$$
 for $0 \le x \le 1$

 $1 + x \le e^x$ for all real x

 $(1-x)^n \ge 1 - nx$ for $0 \le x \le 1$

 $(x+y)^2 \le 2x^2 + 2y^2$

Triangle Inequality $|\mathbf{x} + \mathbf{y}| \le |\mathbf{x}| + |\mathbf{y}|.$

Cauchy-Schwartz Inequality $|\mathbf{x}||\mathbf{y}| \ge \mathbf{x}^T \mathbf{y}$

Young's Inequality For positive real numbers p and q where $\frac{1}{p} + \frac{1}{q} = 1$ and positive reals x and y,

$$xy \le \frac{1}{p}x^p + \frac{1}{q}y^q.$$

Hölder's inequalityHölder's inequality For positive real numbers p and q with $\frac{1}{p} + \frac{1}{q} = 1$,

$$\sum_{i=1}^{n} |x_i y_i| \leq \left(\sum_{i=1}^{n} |x_i|^p\right)^{1/p} \left(\sum_{i=1}^{n} |y_i|^q\right)^{1/q}.$$

Jensen's inequality For a convex function f,

$$f\left(\sum_{i=1}^{n} \alpha_{i} x_{i}\right) \leq \sum_{i=1}^{n} \alpha_{i} f\left(x_{i}\right),$$

Let $g(x) = (1 - x)^n - (1 - nx)$. We establish $g(x) \ge 0$ for x in [0, 1] by taking the derivative.

$$g'(x) = -n(1-x)^{n-1} + n = n\left(1 - (1-x)^{n-1}\right) \ge 0$$

for $0 \le x \le 1$. Thus, g takes on its minimum for x in [0, 1] at x = 0 where g(0) = 0 proving the inequality.

 $(x+y)^2 \leq 2x^2+2y^2$

The inequality follows from $(x + y)^2 + (x - y)^2 = 2x^2 + 2y^2$.

Lemma 12.1 For any nonnegative reals a_1, a_2, \ldots, a_n and any $\rho \in [0, 1]$, $\left(\sum_{i=1}^n a_i\right)^{\rho} \leq \sum_{i=1}^n a_i^{\rho}$.

Proof: We will see that we can reduce the proof of the lemma to the case when only one of the a_i is nonzero and the rest are zero. To this end, suppose a_1 and a_2 are both positive and without loss of generality, assume $a_1 \ge a_2$. Add an infinitesimal positive amount ϵ to a_1 and subtract the same amount from a_2 . This does not alter the left hand side. We claim it does not increase the right hand side. To see this, note that

$$(a_1 + \epsilon)^{\rho} + (a_2 - \epsilon)^{\rho} - a_1^{\rho} - a_2^{\rho} = \rho(a_1^{\rho-1} - a_2^{\rho-1})\epsilon + O(\epsilon^2),$$

and since $\rho - 1 \leq 0$, we have $a_1^{\rho-1} - a_2^{\rho-1} \leq 0$, proving the claim. Now by repeating this process, we can make $a_2 = 0$ (at that time a_1 will equal the sum of the original a_1 and a_2). Now repeating on all pairs of a_i , we can make all but one of them zero and in the process, we have left the left hand side the same, but have not increased the right hand side. So it suffices to prove the inequality at the end which clearly holds. This method of proof is called the variational method.

The Triangle Inequality

For any two vectors \mathbf{x} and \mathbf{y} , $|\mathbf{x} + \mathbf{y}| \le |\mathbf{x}| + |\mathbf{y}|$. Since $\mathbf{x} \cdot \mathbf{y} \le |\mathbf{x}| |\mathbf{y}|$,

 $|\mathbf{x} + \mathbf{y}|^2 = (\mathbf{x} + \mathbf{y})^T \cdot (\mathbf{x} + \mathbf{y}) = |\mathbf{x}|^2 + |\mathbf{y}|^2 + 2\mathbf{x}^T \cdot \mathbf{y} \le |\mathbf{x}|^2 + |\mathbf{y}|^2 + 2|\mathbf{x}||\mathbf{y}| = (|\mathbf{x}| + |\mathbf{y}|)^2.$ The inequality follows by taking square roots.

Stirling approximation

$$n! \cong \left(\frac{n}{e}\right)^n \sqrt{2\pi n} \qquad \qquad \begin{pmatrix} 2n\\n \end{pmatrix} \cong \frac{1}{\sqrt{\pi n}} 2^{2n}$$
$$\sqrt{2\pi n} \frac{n^n}{e^n} < n! < \sqrt{2\pi n} \frac{n^n}{e^n} \left(1 + \frac{1}{12n - 1}\right)$$

We prove the inequalities, except for constant factors. Namely, we prove that

1.4
$$\left(\frac{n}{e}\right)^n \sqrt{n} \le n! \le e \left(\frac{n}{e}\right)^n \sqrt{n}.$$

Write $\ln(n!) = \ln 1 + \ln 2 + \dots + \ln n$. This sum is approximately $\int_{x=1}^{n} \ln x \, dx$. The indefinite integral $\int \ln x \, dx = (x \ln x - x)$ gives an approximation, but without the \sqrt{n} term. To get the \sqrt{n} , differentiate twice and note that $\ln x$ is a concave function. This means that for any positive x_0 ,

$$\frac{\ln x_0 + \ln(x_0 + 1)}{2} \le \int_{x=x_0}^{x_0 + 1} \ln x \, dx,$$

since for $x \in [x_0, x_0 + 1]$, the curve $\ln x$ is always above the spline joining $(x_0, \ln x_0)$ and $(x_0 + 1, \ln(x_0 + 1))$. Thus,

$$\ln(n!) = \frac{\ln 1}{2} + \frac{\ln 1 + \ln 2}{2} + \frac{\ln 2 + \ln 3}{2} + \dots + \frac{\ln(n-1) + \ln n}{2} + \frac{\ln n}{2}$$
$$\leq \int_{x=1}^{n} \ln x \, dx + \frac{\ln n}{2} = [x \ln x - x]_{1}^{n} + \frac{\ln n}{2}$$
$$= n \ln n - n + 1 + \frac{\ln n}{2}.$$

Thus, $n! \leq n^n e^{-n} \sqrt{n} e$. For the lower bound on n!, start with the fact that for any $x_0 \geq 1/2$ and any real ρ

$$\ln x_0 \ge \frac{1}{2} (\ln(x_0 + \rho) + \ln(x_0 - \rho)) \quad \text{implies} \quad \ln x_0 \ge \int_{x=x_0 - 0.5}^{x_0 + .5} \ln x \, dx.$$

Thus,

$$\ln(n!) = \ln 2 + \ln 3 + \dots + \ln n \ge \int_{x=1.5}^{n+.5} \ln x \, dx,$$

from which one can derive a lower bound with a calculation.

Stirling approximation for the binomial coefficient

$$\binom{n}{k} \le \left(\frac{en}{k}\right)^k$$

Using the Stirling approximation for k!,

$$\binom{n}{k} = \frac{n!}{(n-k)!k!} \le \frac{n^k}{k!} \cong \left(\frac{en}{k}\right)^k.$$

The gamma function

For a > 0

$$\Gamma(a) = \int_{0}^{\infty} x^{a-1} e^{-x} dx$$

$$\Gamma\left(\frac{1}{2}\right) = \sqrt{\pi}, \quad \Gamma(1) = \Gamma(2) = 1, \text{ and for } n \ge 2, \quad \Gamma(n) = (n-1)\Gamma(n-1).$$

The last statement is proved by induction on n. It is easy to see that $\Gamma(1) = 1$. For $n \ge 2$, we use integration by parts.

$$\int f(x) g'(x) dx = f(x) g(x) - \int f'(x) g(x) dx$$

Write $\Gamma(n) = \int_{x=0}^{\infty} f(x)g'(x) dx$, where, $f(x) = x^{n-1}$ and $g'(x) = e^{-x}$. Thus,

$$\Gamma(n) = [f(x)g(x)]_{x=0}^{\infty} + \int_{x=0}^{\infty} (n-1)x^{n-2}e^{-x} dx = (n-1)\Gamma(n-1),$$

as claimed.

Cauchy-Schwartz Inequality

$$\left(\sum_{i=1}^{n} x_i^2\right) \left(\sum_{i=1}^{n} y_i^2\right) \ge \left(\sum_{i=1}^{n} x_i y_i\right)^2$$

In vector form, $|\mathbf{x}||\mathbf{y}| \ge \mathbf{x}^T \mathbf{y}$, the inequality states that the dot product of two vectors is at most the product of their lengths. The Cauchy-Schwartz inequality is a special case of Hölder's inequality with p = q = 2.

Young's inequality

For positive real numbers p and q where $\frac{1}{p} + \frac{1}{q} = 1$ and positive reals x and y,

$$\frac{1}{p}x^p + \frac{1}{q}y^q \ge xy.$$

The left hand side of Young's inequality, $\frac{1}{p}x^p + \frac{1}{q}y^q$, is a convex combination of x^p and y^q since $\frac{1}{p}$ and $\frac{1}{q}$ sum to 1. $\ln(x)$ is a concave function for x > 0 and so the ln of the convex combination of the two elements is greater than or equal to the convex combination of the ln of the two elements

$$\ln(\frac{1}{p}x^{p} + \frac{1}{q}y^{p}) \ge \frac{1}{p}\ln(x^{p}) + \frac{1}{q}\ln(y^{q}) = \ln(xy).$$

Since for $x \ge 0$, $\ln x$ is a monotone increasing function, $\frac{1}{p}x^p + \frac{1}{q}y^q \ge xy$.

Hölder's inequalityHölder's inequality

For positive real numbers p and q with $\frac{1}{p} + \frac{1}{q} = 1$,

$$\sum_{i=1}^{n} |x_i y_i| \leq \left(\sum_{i=1}^{n} |x_i|^p \right)^{1/p} \left(\sum_{i=1}^{n} |y_i|^q \right)^{1/q}.$$

Let $x'_i = x_i / (\sum_{i=1}^n |x_i|^p)^{1/p}$ and $y'_i = y_i / (\sum_{i=1}^n |y_i|^q)^{1/q}$. Replacing x_i by x'_i and y_i by y'_i does not change the inequality. Now $\sum_{i=1}^n |x'_i|^p = \sum_{i=1}^n |y'_i|^q = 1$, so it suffices to prove $\sum_{i=1}^n |x'_iy'_i| \le 1$. Apply Young's inequality to get $|x'_iy'_i| \le \frac{|x'_i|^p}{p} + \frac{|y'_i|^q}{q}$. Summing over i, the right hand side sums to $\frac{1}{p} + \frac{1}{q} = 1$ finishing the proof.

For a_1, a_2, \ldots, a_n real and k a positive integer,

$$(a_1 + a_2 + \dots + a_n)^k \le n^{k-1} (|a_1|^k + |a_2|^k + \dots + |a_n|^k).$$

Using Hölder's inequality with p = k and q = k/(k-1),

$$|a_1 + a_2 + \dots + a_n| \le |a_1 \cdot 1| + |a_2 \cdot 1| + \dots + |a_n \cdot 1|$$
$$\le \left(\sum_{i=1}^n |a_i|^k\right)^{1/k} (1 + 1 + \dots + 1)^{(k-1)/k},$$

from which the current inequality follows.

Arithmetic and geometric means

The arithmetic mean of a set of nonnegative reals is at least their geometric mean. For $a_1, a_2, \ldots, a_n > 0$,

$$\frac{1}{n}\sum_{i=1}^{n}a_i \ge \sqrt[n]{a_1a_2\cdots a_n}.$$

Assume that $a_1 \ge a_2 \ge \ldots \ge a_n$. We reduce the proof to the case when all the a_i are equal using the variational method; in this case the inequality holds with equality. Suppose $a_1 > a_2$. Let ε be a positive infinitesimal. Add ε to a_2 and subtract ε from a_1 to get closer to the case when they are equal. The left hand side $\frac{1}{n} \sum_{i=1}^{n} a_i$ does not change.

$$(a_1 - \varepsilon)(a_2 + \varepsilon)a_3a_4 \cdots a_n = a_1a_2 \cdots a_n + \varepsilon(a_1 - a_2)a_3a_4 \cdots a_n + O(\varepsilon^2)$$

> $a_1a_2 \cdots a_n$

for small enough $\varepsilon > 0$. Thus, the change has increased $\sqrt[n]{a_1 a_2 \cdots a_n}$. So if the inequality holds after the change, it must hold before. By continuing this process, one can make all the a_i equal.

Approximating sums by integrals

For monotonic decreasing f(x),

$$\int_{x=m}^{n+1} f(x) dx \le \sum_{i=m}^{n} f(i) \le \int_{x=m-1}^{n} f(x) dx.$$

See Fig. 12.1. Thus,

$$\int_{x=2}^{n+1} \frac{1}{x^2} dx \le \sum_{i=2}^{n} \frac{1}{i^2} = \frac{1}{4} + \frac{1}{9} + \dots + \frac{1}{n^2} \le \int_{x=1}^{n} \frac{1}{x^2} dx$$

and hence $\frac{3}{2} - \frac{1}{n+1} \le \sum_{i=1}^{n} \frac{1}{i^2} \le 2 - \frac{1}{n}$.

Jensen's Inequality

For a convex function f,

$$f\left(\frac{1}{2}(x_1+x_2)\right) \le \frac{1}{2}(f(x_1)+f(x_2)).$$

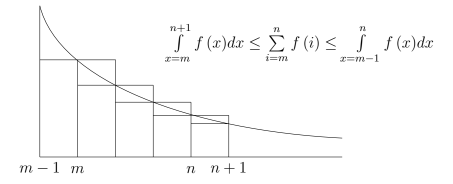


Figure 12.1: Approximating sums by integrals

More generally for any convex function f,

$$f\left(\sum_{i=1}^{n} \alpha_{i} x_{i}\right) \leq \sum_{i=1}^{n} \alpha_{i} f\left(x_{i}\right),$$

where $0 \le \alpha_i \le 1$ and $\sum_{i=1}^n \alpha_i = 1$. From this, it follows that for any convex function f and random variable x,

$$E\left(f\left(x\right)\right) \ge f\left(E\left(x\right)\right).$$

We prove this for a discrete random variable x taking on values a_1, a_2, \ldots with $\operatorname{Prob}(x = a_i) = \alpha_i$:

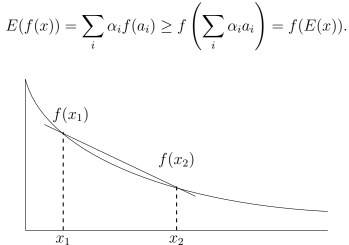


Figure 12.2: For a convex function f, $f\left(\frac{x_1+x_2}{2}\right) \leq \frac{1}{2} \left(f\left(x_1\right) + f\left(x_2\right)\right)$.

Example: Let $f(x) = x^k$ for k an even positive integer. Then, $f''(x) = k(k-1)x^{k-2}$ which since k-2 is even is nonnegative for all x implying that f is convex. Thus,

$$E(x) \le \sqrt[k]{E(x^k)}$$

since $t^{\frac{1}{k}}$ is a monotone function of t, t > 0. It is easy to see that this inequality does not necessarily hold when k is odd; indeed for odd k, x^k is not a convex function.

Tails of Gaussian

For bounding the tails of Gaussian densities, the following inequality is useful. The proof uses a technique useful in many contexts. For t > 0,

$$\int_{x=t}^{\infty} e^{-x^2} dx \le \frac{e^{-t^2}}{2t}.$$

In proof, first write: $\int_{x=t}^{\infty} e^{-x^2} dx \leq \int_{x=t}^{\infty} \frac{x}{t} e^{-x^2} dx$, using the fact that $x \geq t$ in the range of integration. The latter expression is integrable in closed form since $d(e^{-x^2}) = (-2x)e^{-x^2}$ yielding the claimed bound.

A similar technique yields an upper bound on

$$\int_{x=\beta}^{1} (1-x^2)^{\alpha} dx,$$

for $\beta \in [0, 1]$ and $\alpha > 0$. Just use $(1 - x^2)^{\alpha} \leq \frac{x}{\beta}(1 - x^2)^{\alpha}$ over the range and integrate in closed form the last expression.

$$\begin{split} \int_{x=\beta}^{1} (1-x^2)^{\alpha} dx &\leq \int_{x=\beta}^{1} \frac{x}{\beta} (1-x^2)^{\alpha} dx = \left. \frac{-1}{2\beta(\alpha+1)} (1-x^2)^{\alpha+1} \right|_{x=\beta}^{1} \\ &= \frac{(1-\beta^2)^{\alpha+1}}{2\beta(\alpha+1)} \end{split}$$

12.4 Probability

Consider an experiment such as flipping a coin whose outcome is determined by chance. To talk about the outcome of a particular experiment, we introduce the notion of a *ran-dom variable* whose value is the outcome of the experiment. The set of possible outcomes is called the *sample space*. If the sample space is finite, we can assign a probability of occurrence to each outcome. In some situations where the sample space is infinite, we can assign a probability of occurrence. The probability $p(i) = \frac{6}{\pi^2} \frac{1}{i^2}$ for i an integer greater than or equal to one is such an example. The function assigning the probabilities is called

a probability distribution function.

In many situations, a probability distribution function does not exist. For example, for the uniform probability on the interval [0,1], the probability of any specific value is zero. What we can do is define a *probability density function* p(x) such that

$$\operatorname{Prob}(a < x < b) = \int_{a}^{b} p(x) dx$$

If x is a continuous random variable for which a density function exists, then the *cumulative distribution function* f(a) is defined by

$$f(a) = \int_{-\infty}^{a} p(x) dx$$

which gives the probability that $x \leq a$.

12.4.1 Sample Space, Events, Independence

There may be more than one relevant random variable in a situation. For example, if one tosses n coins, there are n random variables, x_1, x_2, \ldots, x_n , taking on values 0 and 1, a 1 for heads and a 0 for tails. The set of possible outcomes, the sample space, is $\{0, 1\}^n$. An *event* is a subset of the sample space. The event of an odd number of heads, consists of all elements of $\{0, 1\}^n$ with an odd number of 1's.

Let A and B be two events. The joint occurrence of the two events is denoted by $(A \wedge B)$. The *conditional probability* of event A given that event B has occurred is denoted by $\operatorname{Prob}(A|B)$ and is given by

$$\operatorname{Prob}(A|B) = \frac{\operatorname{Prob}(A \land B)}{\operatorname{Prob}(B)}.$$

Events A and B are *independent* if the occurrence of one event has no influence on the probability of the other. That is, $\operatorname{Prob}(A|B) = \operatorname{Prob}(A)$ or equivalently, $\operatorname{Prob}(A \wedge B) = \operatorname{Prob}(A)\operatorname{Prob}(B)$. Two random variables x and y are *independent* if for every possible set A of values for x and every possible set B of values for y, the events x in A and y in B are independent.

A collection of n random variables x_1, x_2, \ldots, x_n is *mutually independent* if for all possible sets A_1, A_2, \ldots, A_n of values of x_1, x_2, \ldots, x_n ,

$$\operatorname{Prob}(x_1 \in A_1, x_2 \in A_2, \dots, x_n \in A_n) = \operatorname{Prob}(x_1 \in A_1) \operatorname{Prob}(x_2 \in A_2) \cdots \operatorname{Prob}(x_n \in A_n).$$

If the random variables are discrete, it would suffice to say that for any real numbers a_1, a_2, \ldots, a_n

$$Prob(x_1 = a_1, x_2 = a_2, \dots, x_n = a_n) = Prob(x_1 = a_1)Prob(x_2 = a_2) \cdots Prob(x_n = a_n).$$

Random variables x_1, x_2, \ldots, x_n are pairwise independent if for any a_i and a_j , $i \neq j$, $\operatorname{Prob}(x_i = a_i, x_j = a_j) = \operatorname{Prob}(x_i = a_i)\operatorname{Prob}(x_j = a_j)$. Mutual independence is much stronger than requiring that the variables are pairwise independent. Consider the example of 2-universal hash functions discussed in Chapter ??.

If (x, y) is a random vector and one normalizes it to a unit vector $\left(\frac{x}{\sqrt{x^2+y^2}}, \frac{y}{\sqrt{x^2+y^2}}\right)$ the coordinates are no longer independent since knowing the value of one coordinate uniquely determines the value of the other.

12.4.2 Linearity of Expectation

An important concept is that of the expectation of a random variable. The expected value, E(x), of a random variable x is $E(x) = \sum_{x} xp(x)$ in the discrete case and $E(x) = \int_{-\infty}^{\infty} xp(x)dx$ in the continuous case. The expectation of a sum of random variables is equal to the sum of their expectations. The linearity of expectation follows directly from the definition and does not require independence.

12.4.3 Union Bound

Let A_1, A_2, \ldots, A_n be events. The actual probability of the union of events is given by Boole's formula.

$$\operatorname{Prob}(A_1 \cup A_2 \cup \cdots \cup A_n) = \sum_{i=1}^n \operatorname{Prob}(A_i) - \sum_{ij} \operatorname{Prob}(A_i \wedge A_j) + \sum_{ijk} \operatorname{Prob}(A_i \wedge A_j \wedge A_k) - \cdots$$

Often we only need an upper bound on the probability of the union and use

$$\operatorname{Prob}(A_1 \cup A_2 \cup \cdots \cap A_n) \leq \sum_{i=1}^n \operatorname{Prob}(A_i)$$

This upper bound is called the *union bound*.

12.4.4 Indicator Variables

A useful tool is that of an indicator variable that takes on value 0 or 1 to indicate whether some quantity is present or not. The indicator variable is useful in determining the expected size of a subset. Given a random subset of the integers $\{1, 2, ..., n\}$, the expected size of the subset is the expected value of $x_1 + x_2 + \cdots + x_n$ where x_i is the indicator variable that takes on value 1 if *i* is in the subset.

Example: Consider a random permutation of n integers. Define the indicator function $x_i = 1$ if the i^{th} integer in the permutation is i. The expected number of fixed points is given by

$$E\left(\sum_{i=1}^{n} x_i\right) = \sum_{i=1}^{n} E(x_i) = n\frac{1}{n} = 1.$$

Note that the x_i are not independent. But, linearity of expectation still applies.

Example: Consider the expected number of vertices of degree d in a random graph G(n, p). The number of vertices of degree d is the sum of n indicator random variables, one for each vertex, with value one if the vertex has degree d. The expectation is the sum of the expectations of the n indicator random variables and this is just n times the expectation of one of them. Thus, the expected number of degree d vertices is $n \binom{n}{d} p^d (1-p)^{n-d}$.

12.4.5 Variance

In addition to the expected value of a random variable, another important parameter is the variance. The variance of a random variable x, denoted $\operatorname{var}(x)$ or often $\sigma^2(x)$ is $E(x - E(x))^2$ and measures how close to the expected value the random variable is likely to be. The standard deviation σ is the square root of the variance. The units of σ are the same as those of x.

By linearity of expectation

$$\sigma^{2} = E \left(x - E \left(x \right) \right)^{2} = E(x^{2}) - 2E(x)E(x) + E^{2}(x) = E \left(x^{2} \right) - E^{2}(x) \,.$$

12.4.6 Variance of the Sum of Independent Random Variables

In general, the variance of the sum is not equal to the sum of the variances. However, if x and y are independent, then E(xy) = E(x) E(y) and

$$var(x+y) = var(x) + var(y).$$

To see this

$$var(x+y) = E((x+y)^2) - E^2(x+y)$$

= $E(x^2) + 2E(xy) + E(y^2) - E^2(x) - 2E(x)E(y) - E^2(y).$

From independence, 2E(xy) - 2E(x)E(y) = 0 and

$$var(x + y) = E(x^2) - E^2(x) + E(y^2) - E^2(y)$$

= $var(x) + var(y)$.

More generally, if x_1, x_2, \ldots, x_n are pairwise independent random variables, then

$$var(x_1 + x_2 + \dots + x_n) = var(x_1) + var(x_2) + \dots + var(x_n)$$

For the variance of the sum to be the sum of the variances only requires pairwise independence not full independence.

12.4.7 Median

One often calculates the average value of a random variable to get a feeling for the magnitude of the variable. This is reasonable when the probability distribution of the variable is Gaussian, or has a small variance. However, if there are outliers, then the average may be distorted by outliers. An alternative to calculating the expected value is to calculate the median, the value for which half of the probability is above and half is below.

12.4.8 The Central Limit Theorem

Let $s = x_1 + x_2 + \cdots + x_n$ be a sum of *n* independent random variables where each x_i has probability distribution

$$x_i = \begin{cases} 0 & \frac{1}{2} \\ 1 & \frac{1}{2} \end{cases}$$

The expected value of each x_i is $\frac{1}{2}$ with variance

$$\sigma_i^2 = \left(\frac{1}{2} - 0\right)^2 \frac{1}{2} + \left(\frac{1}{2} - 1\right)^2 \frac{1}{2} = \frac{1}{4}.$$

The expected value of s is n/2 and since the variables are independent, the variance of the sum is the sum of the variances and hence is n/4. How concentrated s is around its mean depends on the standard deviation of s which is $\frac{\sqrt{n}}{2}$. For n equal 100 the expected value of s is 50 with a standard deviation of 5 which is 10% of the mean. For n = 10,000 the expected value of s is 5,000 with a standard deviation of 50 which is 1% of the mean. Note that as n increases, the standard deviation increases, but the ratio of the standard deviation to the mean goes to zero. More generally, if x_i are independent and identically distributed, each with standard deviation σ , then the standard deviation of $x_1 + x_2 + \cdots + x_n$ is $\sqrt{n\sigma}$. So, $\frac{x_1 + x_2 + \cdots + x_n}{\sqrt{n}}$ has standard deviation σ . The central limit theorem makes a stronger assertion that in fact $\frac{x_1 + x_2 + \cdots + x_n}{\sqrt{n}}$ has Gaussian distribution with standard deviation σ .

Theorem 12.2 Suppose x_1, x_2, \ldots, x_n is a sequence of identically distributed independent random variables, each with mean μ and variance σ^2 . The distribution of the random variable

$$\frac{1}{\sqrt{n}}\left(x_1 + x_2 + \dots + x_n - n\mu\right)$$

converges to the distribution of the Gaussian with mean 0 and variance σ^2 .

12.4.9 Probability Distributions

The Gaussian or normal distribution

The normal distribution is

$$\frac{1}{\sqrt{2\pi}\sigma}e^{-\frac{1}{2}\frac{(x-m)^2}{\sigma^2}}$$

where m is the mean and σ^2 is the variance. The coefficient $\frac{1}{\sqrt{2\pi\sigma}}$ makes the integral of the distribution be one. If we measure distance in units of the standard deviation σ from the mean, then

$$\phi(x) = \frac{1}{\sqrt{2\pi}} e^{-\frac{1}{2}x^2}$$

Standard tables give values of the integral

$$\int_{0}^{t} \phi(x) dx$$

and from these values one can compute probability integrals for a normal distribution with mean m and variance σ^2 .

General Gaussians

So far we have seen spherical Gaussian densities in \mathbf{R}^d . The word spherical indicates that the level curves of the density are spheres. If a random vector \mathbf{y} in \mathbf{R}^d has a spherical Gaussian density with zero mean, then y_i and y_j , $i \neq j$, are independent. However, in many situations the variables are correlated. To model these Gaussians, level curves that are ellipsoids rather than spheres are used.

For a random vector \mathbf{x} , the covariance of x_i and x_j is $E((x_i - \mu_i)(x_j - \mu_j))$. We list the covariances in a matrix called the *covariance matrix*, denoted Σ .³⁶ Since \mathbf{x} and $\boldsymbol{\mu}$ are column vectors, $(\mathbf{x} - \boldsymbol{\mu})(\mathbf{x} - \boldsymbol{\mu})^T$ is a $d \times d$ matrix. Expectation of a matrix or vector means componentwise expectation.

$$\Sigma = E((\mathbf{x} - \boldsymbol{\mu})(\mathbf{x} - \boldsymbol{\mu})^T).$$

The general Gaussian density with mean μ and positive definite covariance matrix Σ is

$$f(\mathbf{x}) = \frac{1}{\sqrt{(2\pi)^d \det(\Sigma)}} \exp\left(-\frac{1}{2}(\mathbf{x} - \boldsymbol{\mu})^T \Sigma^{-1}(\mathbf{x} - \boldsymbol{\mu})\right).$$

To compute the covariance matrix of the Gaussian, substitute $\mathbf{y} = \Sigma^{-1/2} (\mathbf{x} - \boldsymbol{\mu})$. Noting that a positive definite symmetric matrix has a square root:

$$E((\mathbf{x} - \boldsymbol{\mu})(\mathbf{x} - \boldsymbol{\mu})^T = E(\Sigma^{1/2}\mathbf{y}\mathbf{y}^T\Sigma^{1/2})$$
$$= \Sigma^{1/2} \left(E(\mathbf{y}\mathbf{y}^T)\right)\Sigma^{1/2} = \Sigma$$

 $^{^{36}\}Sigma$ is the standard notation for the covariance matrix. We will use it sparingly so as not to confuse with the summation sign.

The density of **y** is the unit variance, zero mean Gaussian, thus $E(yy^T) = I$.

Bernoulli trials and the binomial distribution

A Bernoulli trial has two possible outcomes, called success or failure, with probabilities p and 1 - p, respectively. If there are n independent Bernoulli trials, the probability of exactly k successes is given by the *binomial distribution*

$$B(n,p) = \binom{n}{k} p^k (1-p)^{n-k}$$

The mean and variance of the binomial distribution B(n, p) are np and np(1-p), respectively. The mean of the binomial distribution is np, by linearity of expectations. The variance is np(1-p) since the variance of a sum of independent random variables is the sum of their variances.

Let x_1 be the number of successes in n_1 trials and let x_2 be the number of successes in n_2 trials. The probability distribution of the sum of the successes, $x_1 + x_2$, is the same as the distribution of $x_1 + x_2$ successes in $n_1 + n_2$ trials. Thus, $B(n_1, p) + B(n_2, p) = B(n_1 + n_2, p)$.

When p is a constant, the expected degree of vertices in G(n, p) increases with n. For example, in $G(n, \frac{1}{2})$, the expected degree of a vertex is n/2. In many real applications, we will be concerned with G(n, p) where p = d/n, for d a constant; i.e., graphs whose expected degree is a constant d independent of n. Holding d = np constant as n goes to infinity, the binomial distribution

$$\operatorname{Prob}\left(k\right) = \binom{n}{k} p^{k} \left(1-p\right)^{n-k}$$

approaches the Poisson distribution

$$Prob(k) = \frac{(np)^k}{k!}e^{-np} = \frac{d^k}{k!}e^{-d}.$$

To see this, assume k = o(n) and use the approximations $n - k \cong n$, $\binom{n}{k} \cong \frac{n^k}{k!}$, and $\left(1 - \frac{1}{n}\right)^{n-k} \cong e^{-1}$ to approximate the binomial distribution by

$$\lim_{n \to \infty} \binom{n}{k} p^k (1-p)^{n-k} = \frac{n^k}{k!} \left(\frac{d}{n}\right)^k (1-\frac{d}{n})^n = \frac{d^k}{k!} e^{-d}.$$

Note that for $p = \frac{d}{n}$, where d is a constant independent of n, the probability of the binomial distribution falls off rapidly for k > d, and is essentially zero for all but some finite number of values of k. This justifies the k = o(n) assumption. Thus, the Poisson distribution is a good approximation.

Poisson distribution

The Poisson distribution describes the probability of k events happening in a unit of time when the average rate per unit of time is λ . Divide the unit of time into n segments. When n is large enough, each segment is sufficiently small so that the probability of two events happening in the same segment is negligible. The Poisson distribution gives the probability of k events happening in a unit of time and can be derived from the binomial distribution by taking the limit as $n \to \infty$.

Let
$$p = \frac{\lambda}{n}$$
. Then
Prob $(k \text{ successes in a unit of time}) = \lim_{n \to \infty} {\binom{n}{k} \left(\frac{\lambda}{n}\right)^k \left(1 - \frac{\lambda}{n}\right)^{n-k}}$
 $= \lim_{n \to \infty} \frac{n \left(n - 1\right) \cdots \left(n - k + 1\right)}{k!} \left(\frac{\lambda}{n}\right)^k \left(1 - \frac{\lambda}{n}\right)^n \left(1 - \frac{\lambda}{n}\right)^{-k}$
 $= \lim_{n \to \infty} \frac{\lambda^k}{k!} e^{-\lambda}$

In the limit as n goes to infinity the binomial distribution $p(k) = {n \choose k} p^k (1-p)^{n-k}$ becomes the Poisson distribution $p(k) = e^{-\lambda} \frac{\lambda^k}{k!}$. The mean and the variance of the Poisson distribution have value λ . If x and y are both Poisson random variables from distributions with means λ_1 and λ_2 respectively, then x + y is Poisson with mean $m_1 + m_2$. For large n and small p the binomial distribution can be approximated with the Poisson distribution.

The binomial distribution with mean np and variance np(1-p) can be approximated by the normal distribution with mean np and variance np(1-p). The central limit theorem tells us that there is such an approximation in the limit. The approximation is good if both np and n(1-p) are greater than 10 provided k is not extreme. Thus,

$$\binom{n}{k} \left(\frac{1}{2}\right)^k \left(\frac{1}{2}\right)^{n-k} \cong \frac{1}{\sqrt{\pi n/2}} e^{-\frac{(n/2-k)^2}{\frac{1}{2}n}}$$

This approximation is excellent provided k is $\Theta(n)$. The Poisson approximation

$$\binom{n}{k} p^k \left(1-p\right)^k \cong e^{-np} \frac{\left(np\right)^k}{k!}$$

is off for central values and tail values even for p = 1/2. The approximation

$$\binom{n}{k} p^k (1-p)^{n-k} \cong \frac{1}{\sqrt{\pi pn}} e^{-\frac{(pn-k)^2}{pn}}$$

is good for p = 1/2 but is off for other values of p.

Generation of random numbers according to a given probability distribution

Suppose one wanted to generate a random variable with probability density p(x) where p(x) is continuous. Let P(x) be the cumulative distribution function for x and let u be a random variable with uniform probability density over the interval [0,1]. Then the random variable $x = P^{-1}(u)$ has probability density p(x).

Example: For a Cauchy density function the cumulative distribution function is

$$P(x) = \int_{t=-\infty}^{x} \frac{1}{\pi} \frac{1}{1+t^2} dt = \frac{1}{2} + \frac{1}{\pi} \tan^{-1}(x).$$

Setting u = P(x) and solving for x yields $x = \tan\left(\pi\left(u - \frac{1}{2}\right)\right)$. Thus, to generate a random number $x \ge 0$ using the Cauchy distribution, generate $u, 0 \le u \le 1$, uniformly and calculate $x = \tan\left(\pi\left(u - \frac{1}{2}\right)\right)$. The value of x varies from $-\infty$ to ∞ with x = 0 for u = 1/2.

12.4.10 Bayes Rule and Estimators

Bayes rule

Bayes rule relates the conditional probability of A given B to the conditional probability of B given A.

$$\operatorname{Prob}\left(A|B\right) = \frac{\operatorname{Prob}\left(B|A\right)\operatorname{Prob}\left(A\right)}{\operatorname{Prob}\left(B\right)}$$

Suppose one knows the probability of A and wants to know how this probability changes if we know that B has occurred. Prob(A) is called the prior probability. The conditional probability Prob(A|B) is called the posterior probability because it is the probability of A after we know that B has occurred.

The example below illustrates that if a situation is rare, a highly accurate test will often give the wrong answer.

Example: Let A be the event that a product is defective and let B be the event that a test says a product is defective. Let $\operatorname{Prob}(B|A)$ be the probability that the test says a product is defective assuming the product is defective and let $\operatorname{Prob}(B|\overline{A})$ be the probability that the test says a product is defective if it is not actually defective.

What is the probability $\operatorname{Prob}(A|B)$ that the product is defective if the test say it is defective? Suppose $\operatorname{Prob}(A) = 0.001$, $\operatorname{Prob}(B|A) = 0.99$, and $\operatorname{Prob}(B|\overline{A}) = 0.02$. Then

$$Prob (B) = Prob (B|A) Prob (A) + Prob (B|\overline{A}) Prob (\overline{A})$$
$$= 0.99 \times 0.001 + 0.02 \times 0.999$$
$$= 0.02087$$

and

$$\operatorname{Prob}(A|B) = \frac{\operatorname{Prob}(B|A)\operatorname{Prob}(A)}{\operatorname{Prob}(B)} \approx \frac{0.99 \times 0.001}{0.0210} = 0.0471$$

Even though the test fails to detect a defective product only 1% of the time when it is defective and claims that it is defective when it is not only 2% of the time, the test is correct only 4.7% of the time when it says a product is defective. This comes about because of the low frequencies of defective products.

The words prior, a posteriori, and likelihood come from Bayes theorem.

a posteriori =
$$\frac{\text{likelihood} \times \text{prior}}{\text{normalizing constant}}$$

 $\text{Prob}(A|B) = \frac{\text{Prob}(B|A) \text{Prob}(A)}{\text{Prob}(B)}$

The a posteriori probability is the conditional probability of A given B. The likelihood is the conditional probability $\operatorname{Prob}(B|A)$.

Unbiased Estimators

Consider *n* samples x_1, x_2, \ldots, x_n from a Gaussian distribution of mean μ and variance σ^2 . For this distribution, $m = \frac{x_1 + x_2 + \cdots + x_n}{n}$ is an unbiased estimator of μ , which means that $E(m) = \mu$ and $\frac{1}{n} \sum_{i=1}^n (x_i - \mu)^2$ is an unbiased estimator of σ^2 . However, if μ is not known and is approximated by *m*, then $\frac{1}{n-1} \sum_{i=1}^n (x_i - m)^2$ is an unbiased estimator of σ^2 .

Maximum Likelihood Estimation MLE

Suppose the probability distribution of a random variable x depends on a parameter r. With slight abuse of notation, since r is a parameter rather than a random variable, we denote the probability distribution of x as p(x|r). This is the likelihood of observing x if r was in fact the parameter value. The job of the maximum likelihood estimator, MLE, is to find the best r after observing values of the random variable x. The likelihood of r being the parameter value given that we have observed x is denoted L(r|x). This is again not a probability since r is a parameter, not a random variable. However, if we were to apply Bayes' rule as if this was a conditional probability, we get

$$L(r|x) = \frac{\operatorname{Prob}(x|r)\operatorname{Prob}(r)}{\operatorname{Prob}(x)}$$

Now, assume $\operatorname{Prob}(r)$ is the same for all r. The denominator $\operatorname{Prob}(x)$ is the absolute probability of observing x and is independent of r. So to maximize L(r|x), we just maximize $\operatorname{Prob}(x|r)$. In some situations, one has a prior guess as to the distribution $\operatorname{Prob}(r)$. This is then called the "prior" and in that case, we call $\operatorname{Prob}(x|r)$ the posterior which we try to maximize.

Example: Consider flipping a coin 100 times. Suppose 62 heads and 38 tails occur. What is the most likely value of the probability of the coin to come down heads when the coin is flipped? In this case, it is r = 0.62. The probability that we get 62 heads if the unknown probability of heads in one trial is r is

Prob (62 heads|r) =
$$\binom{100}{62} r^{62} (1-r)^{38}$$
.

This quantity is maximized when r = 0.62. To see this take the logarithm, which as a function of r is $\ln \binom{100}{62} + 62 \ln r + 38 \ln(1-r)$. The derivative with respect to r is zero at r = 0.62 and the second derivative is negative indicating a maximum. Thus, r = 0.62 is the maximum likelihood estimator of the probability of heads in a trial.

12.4.11 Tail Bounds and Chernoff inequalities

Markov's inequality bounds the probability that a nonnegative random variable exceeds a value a. E(r)

or

$$p(x \ge a) \le \frac{D(w)}{a}.$$
$$p(x \ge aE(x)) \le \frac{1}{a}.$$

If one also knows the variance, σ^2 , then using Chebyshev's inequality one can bound the probability that a random variable differs from its expected value by more than *a* standard deviations.

$$p(|x-m| \ge a\sigma) \le \frac{1}{a^2}$$

If a random variable s is the sum of n independent random variables x_1, x_2, \ldots, x_n of finite variance, then better bounds are possible. For any $\delta > 0$,

$$\operatorname{Prob}(s > (1+\delta)m) < \left[\frac{e^{\delta}}{(1+\delta)^{(1+\delta)}}\right]^m$$

and for $0 < \gamma \leq 1$,

Prob
$$(s < (1 - \gamma)m) < \left[\frac{e^{-\gamma}}{(1 + \gamma)^{(1+\gamma)}}\right]^m < e^{-\frac{\gamma^2 m}{2}}$$

Chernoff inequalities

Chebyshev's inequality bounds the probability that a random variable will deviate from its mean by more than a given amount. Chebyshev's inequality holds for any probability distribution. For some distributions we can get much tighter bounds. For example, the probability that a Gaussian random variable deviates from its mean falls off exponentially with the distance from the mean. Here we shall be concerned with the situation where we have a random variable that is the sum of n independent random variables. This is another situation in which we can derive a tighter bound than that given by the Chebyshev inequality. We consider the case where the n independent variables are binomial but similar results can be shown for independent random variables from any distribution that has a finite variance.

Let x_1, x_2, \ldots, x_n be independent random variables where

$$x_i = \begin{cases} 0 & \operatorname{Prob} \ 1 - p \\ 1 & \operatorname{Prob} \ p \end{cases}$$

.

Consider the sum $s = \sum_{i=1}^{n} x_i$. Here the expected value of each x_i is p and by linearity of expectation, the expected value of the sum is m=np. Theorem 12.5 bounds the probability that the sum s exceeds $(1 + \delta) m$.

Theorem 12.3 For any $\delta > 0$, $Prob\left(s > (1+\delta)m\right) < \left(\frac{e^{\delta}}{(1+\delta)^{(1+\delta)}}\right)^m$

Proof: For any $\lambda > 0$, the function $e^{\lambda x}$ is monotone. Thus,

$$\operatorname{Prob}\left(s > (1+\delta)m\right) = \operatorname{Prob}\left(e^{\lambda s} > e^{\lambda(1+\delta)m}\right).$$

 $e^{\lambda x}$ is nonnegative for all x, so we can apply Markov's inequality to get

Prob
$$\left(e^{\lambda s} > e^{\lambda(1+\delta)m}\right) \leq e^{-\lambda(1+\delta)m} E\left(e^{\lambda s}\right).$$

Since the x_i are independent,

$$E\left(e^{\lambda s}\right) = E\left(e^{\lambda \sum_{i=1}^{n} x_{i}}\right) = E\left(\prod_{i=1}^{n} e^{\lambda x_{i}}\right) = \prod_{i=1}^{n} E\left(e^{\lambda x_{i}}\right)$$
$$= \prod_{i=1}^{n} \left(e^{\lambda}p + 1 - p\right) = \prod_{i=1}^{n} \left(p(e^{\lambda} - 1) + 1\right).$$

Using the inequality $1 + x < e^x$ with $x = p(e^{\lambda} - 1)$ yields

$$E\left(e^{\lambda s}\right) < \prod_{i=1}^{n} e^{p(e^{\lambda}-1)}.$$

Thus, for all $\lambda > 0$

$$\operatorname{Prob}\left(s > (1+\delta)m\right) \leq \operatorname{Prob}\left(e^{\lambda s} > e^{\lambda(1+\delta)m}\right)$$
$$\leq e^{-\lambda(1+\delta)m} E\left(e^{\lambda s}\right)$$
$$\leq e^{-\lambda(1+\delta)m} \prod_{i=1}^{n} e^{p(e^{\lambda}-1)}.$$

Setting $\lambda = \ln(1 + \delta)$

$$\operatorname{Prob}\left(s > (1+\delta)m\right) \leq \left(e^{-\ln(1+\delta)}\right)^{(1+\delta)m} \prod_{i=1}^{n} e^{p(e^{\ln(1+\delta)}-1)}$$
$$\leq \left(\frac{1}{1+\delta}\right)^{(1+\delta)m} \prod_{i=1}^{n} e^{p\delta}$$
$$\leq \left(\frac{1}{(1+\delta)}\right)^{(1+\delta)m} e^{np\delta}$$
$$\leq \left(\frac{e^{\delta}}{(1+\delta)^{(1+\delta)}}\right)^{m}.$$

To simplify the bound of Theorem 12.3, observe that

$$(1+\delta)\ln(1+\delta) = \delta + \frac{\delta^2}{2} - \frac{\delta^3}{6} + \frac{\delta^4}{12} - \cdots$$

Therefore

$$(1+\delta)^{(1+\delta)} = e^{\delta + \frac{\delta^2}{2} - \frac{\delta^3}{6} + \frac{\delta^4}{12} - \dots}$$

and hence

$$\frac{e^{\delta}}{(1+\delta)^{(1+\delta)}} = e^{-\frac{\delta^2}{2} + \frac{\delta^3}{6} - \cdots}.$$

Thus, the bound simplifies to

$$\operatorname{Prob}\left(s < (1+\delta)\,m\right) \le e^{-\frac{\delta^2}{2}m + \frac{\delta^3}{6}m - \cdots}.$$

For small δ the probability drops exponentially with δ^2 .

When δ is large another simplification is possible. First

$$\operatorname{Prob}\left(s > (1+\delta) \, m\right) \le \left(\frac{e^{\delta}}{\left(1+\delta\right)^{(1+\delta)}}\right)^m \le \left(\frac{e}{1+\delta}\right)^{(1+\delta)m}$$

If $\delta > 2e - 1$, substituting 2e - 1 for δ in the denominator yields

$$\operatorname{Prob}(s > (1+\delta)m) \le 2^{-(1+\delta)m}$$

Theorem 12.3 gives a bound on the probability of the sum being greater than the mean. We now bound the probability that the sum will be less than its mean.

Theorem 12.4 Let $0 < \gamma \leq 1$, then $\Pr{ob}\left(s < (1-\gamma)m\right) < \left(\frac{e^{-\gamma}}{(1+\gamma)^{(1+\gamma)}}\right)^m < e^{-\frac{\gamma^2 m}{2}}$. **Proof:** For any $\lambda > 0$

 $\operatorname{Prob}\left(s < (1-\gamma)m\right) = \operatorname{Prob}\left(-s > -(1-\gamma)m\right) = \operatorname{Prob}\left(e^{-\lambda s} > e^{-\lambda(1-\gamma)m}\right).$

Applying Markov's inequality

$$\operatorname{Prob}\left(s < (1-\gamma)m\right) < \frac{E(e^{-\lambda x})}{e^{-\lambda(1-\gamma)m}} < \frac{\prod_{i=1}^{n} E(e^{-\lambda X_i})}{e^{-\lambda(1-\gamma)m}}.$$

Now

$$E(e^{-\lambda x_i}) = pe^{-\lambda} + 1 - p = 1 + p(e^{-\lambda} - 1) + 1.$$

Thus,

Prob
$$(s < (1 - \gamma)m) < \frac{\prod_{i=1}^{n} [1 + p(e^{-\lambda} - 1)]}{e^{-\lambda(1 - \gamma)m}}.$$

Since $1 + x < e^x$

Prob
$$\left(s < (1 - \gamma)m\right) < \frac{e^{np(e^{-\lambda} - 1)}}{e^{-\lambda(1 - \gamma)m}}.$$

Setting $\lambda = \ln \frac{1}{1-\gamma}$

$$\operatorname{Prob}\left(s < (1-\gamma)m\right) < \frac{e^{np(1-\gamma-1)}}{(1-\gamma)^{(1-\gamma)m}} < \left(\frac{e^{-\gamma}}{(1-\gamma)^{(1-\gamma)}}\right)^m.$$

But for $0 < \gamma \le 1$, $(1 - \gamma)^{(1-\gamma)} > e^{-\gamma + \frac{\gamma^2}{2}}$. To see this note that

$$(1-\gamma)\ln(1-\gamma) = (1-\gamma)\left(-\gamma - \frac{\gamma^2}{2} - \frac{\gamma^3}{3} - \cdots\right)$$
$$= -\gamma - \frac{\gamma^2}{2} - \frac{\gamma^3}{3} - \cdots + \gamma^2 + \frac{\gamma^3}{2} + \frac{\gamma^4}{3} + \cdots$$
$$= -\gamma + \left(\gamma^2 - \frac{\gamma^2}{2}\right) + \left(\frac{\gamma^3}{2} - \frac{\gamma^3}{3}\right) + \cdots$$
$$= -\gamma + \frac{\gamma^2}{2} + \frac{\gamma^3}{6} + \cdots$$
$$\ge -\gamma + \frac{\gamma^2}{2}.$$

It then follows that

Prob
$$(s < (1 - \gamma)m) < \left(\frac{e^{-\gamma}}{(1 - \gamma)^{(1 - \gamma)}}\right)^m < e^{-\frac{m\gamma^2}{2}}.$$

12.5 Bounds on Tail Probability

After an introduction to tail inequalties, the main purpose of this section is to state the Master Tail bounds theorem of Chapter 2 (with more detail), give a proof of it, and derive the other tail inequalities mentioned in the table in that chapter.

Markov's inequality bounds the tail probability of a nonnegative random variable x based only on its expectation. For a > 0,

$$\Pr(x > a) \le \frac{E(x)}{a}.$$

As a grows, the bound drops off as 1/a. Given the second moment of x, Chebyshev's inequality, which does not assume x is a nonnegative random variable, gives a tail bound falling off as $1/a^2$

$$\Pr(|x - E(x)| \ge a) \le \frac{E\left(\left(x - E(x)\right)^2\right)}{a^2}.$$

Higher moments yield bounds by applying either of these two theorems. For example, if r is a nonnegative even integer, then x^r is a nonnegative random variable even if x takes on negative values. Applying Markov's inequality to x^r ,

$$\Pr(|x| \ge a) = \Pr(x^r \ge a^r) \le \frac{E(x^r)}{a^r},$$

a bound that falls off as $1/a^r$. The larger the r, the greater the rate of fall, but a bound on $E(x^r)$ is needed to apply this technique.

For a random variable x that is the sum of a large number of independent random variables, x_1, x_2, \ldots, x_n , one can derive bounds on $E(x^r)$ for high even r. There are many situations where the sum of a large number of independent random variables arises. For example, x_i may be the amount of a good that the i^{th} consumer buys, the length of the i^{th} message sent over a network, or the indicator random variable of whether the i^{th} record in a large database has a certain property. Each x_i is modeled by a simple probability distribution. Gaussian, exponential probability density (at any t > 0 is e^{-t}), or binomial distributions are typically used, in fact, respectively in the three examples here. If the x_i have 0-1 distributions, there are a number of theorems called Chernoff bounds, bounding the tails of $x = x_1 + x_2 + \cdots + x_n$, typically proved by the so-called moment-generating function method (see Section 12.4.11 of the appendix). But exponential and Gaussian random variables are not bounded and these methods do not apply. However, good bounds on the moments of these two distributions are known. Indeed, for any integer s > 0, the s^{th} moment for the unit variance Gaussian and the exponential are both at most s!.

Given bounds on the moments of individual x_i the following theorem proves moment bounds on their sum. We use this theorem to derive tail bounds not only for sums of 0-1 random variables, but also Gaussians, exponentials, Poisson, etc.

The gold standard for tail bounds is the central limit theorem for independent, identically distributed random variables x_1, x_2, \dots, x_n with zero mean and $\operatorname{Var}(x_i) = \sigma^2$ that states as $n \to \infty$ the distribution of $x = (x_1 + x_2 + \dots + x_n)/\sqrt{n}$ tends to the Gaussian density with zero mean and variance σ^2 . Loosely, this says that in the limit, the tails of $x = (x_1 + x_2 + \dots + x_n)/\sqrt{n}$ are bounded by that of a Gaussian with variance σ^2 . But this theorem is only in the limit, whereas, we prove a bound that applies for all n.

In the following theorem, x is the sum of n independent, not necessarily identically distributed, random variables x_1, x_2, \ldots, x_n , each of zero mean and variance at most σ^2 . By the central limit theorem, in the limit the probability density of x goes to that of the Gaussian with variance at most $n\sigma^2$. In a limit sense, this implies an upper bound of $ce^{-a^2/(2n\sigma^2)}$ for the tail probability $\Pr(|x| > a)$ for some constant c. The following theorem assumes bounds on higher moments, and asserts a quantitative upper bound of $3e^{-a^2/(12n\sigma^2)}$ on the tail probability, not just in the limit, but for every n. We will apply this theorem to get tail bounds on sums of Gaussian, binomial, and power law distributed random variables.

Theorem 12.5 Let $x = x_1 + x_2 + \cdots + x_n$, where x_1, x_2, \ldots, x_n are mutually independent random variables with zero mean and variance at most σ^2 . Suppose $a \in [0, \sqrt{2n\sigma^2}]$ and $s \leq n\sigma^2/2$ is a positive even integer and $|E(x_i^r)| \leq \sigma^2 r!$, for $r = 3, 4, \ldots, s$. Then,

$$Pr(|x_1 + x_2 + \cdots + x_n| \ge a) \le \left(\frac{2sn\sigma^2}{a^2}\right)^{s/2}.$$

If further, $s \ge a^2/(4n\sigma^2)$, then we also have:

$$Pr(|x_1 + x_2 + \dots + x_n| \ge a) \le 3e^{-a^2/(12n\sigma^2)}.$$

Proof: We first prove an upper bound on $E(x^r)$ for any even positive integer r and then use Markov's inequality as discussed earlier. Expand $(x_1 + x_2 + \cdots + x_n)^r$.

$$(x_1 + x_2 + \dots + x_n)^r = \sum {\binom{r}{r_1, r_2, \dots, r_n}} x_1^{r_1} x_2^{r_2} \cdots x_n^{r_n}$$
$$= \sum \frac{r!}{r_1! r_2! \cdots r_n!} x_1^{r_1} x_2^{r_2} \cdots x_n^{r_n}$$

where the r_i range over all nonnegative integers summing to r. By independence

$$E(x^{r}) = \sum \frac{r!}{r_{1}!r_{2}!\cdots r_{n}!} E(x_{1}^{r_{1}})E(x_{2}^{r_{2}})\cdots E(x_{n}^{r_{n}}).$$

If in a term, any $r_i = 1$, the term is zero since $E(x_i) = 0$. Assume henceforth that (r_1, r_2, \ldots, r_n) runs over sets of nonzero r_i summing to r where each nonzero r_i is at least two. There are at most r/2 nonzero r_i in each set. Since $|E(x_i^{r_i})| \leq \sigma^2 r_i!$,

$$E(x^r) \le r! \sum_{(r_1, r_2, \dots, r_n)} \sigma^{2(\text{ number of nonzero } r_i \text{ in set})}.$$

Collect terms of the summation with t nonzero r_i for t = 1, 2, ..., r/2. There are $\binom{n}{t}$ subsets of $\{1, 2, ..., n\}$ of cardinality t. Once a subset is fixed as the set of t values of i with nonzero r_i , set each of the $r_i \ge 2$. That is, allocate two to each of the r_i and then allocate the remaining r - 2t to the $t r_i$ arbitrarily. The number of such allocations is just $\binom{r-2t+t-1}{t-1} = \binom{r-t-1}{t-1}$. So,

$$E(x^r) \le r! \sum_{t=1}^{r/2} f(t)$$
, where $f(t) = \binom{n}{t} \binom{r-t-1}{t-1} \sigma^{2t}$.

Thus $f(t) \le h(t)$, where $h(t) = \frac{(n\sigma^2)^t}{t!} 2^{r-t-1}$. Since $t \le r/2 \le n\sigma^2/4$, we have

$$\frac{h(t)}{h(t-1)} = \frac{n\sigma^2}{2t} \ge 2.$$

So, we get

$$E(x^{r}) = r! \sum_{t=1}^{r/2} f(t) \le r! h(r/2) (1 + \frac{1}{2} + \frac{1}{4} + \dots) \le \frac{r!}{(r/2)!} 2^{r/2} (n\sigma^{2})^{r/2}.$$

Applying Markov inequality,

$$\Pr(|x| > a) = \Pr(|x|^r > a^r) \le \frac{r!(n\sigma^2)^{r/2}2^{r/2}}{(r/2)!a^r} = g(r) \le \left(\frac{2rn\sigma^2}{a^2}\right)^{r/2}$$

This holds for all $r \leq s$, r even and applying it with r = s, we get the first inequality of the theorem.

We now prove the second inequality. For even r, $g(r)/g(r-2) = \frac{4(r-1)n\sigma^2}{a^2}$ and so g(r) decreases as long as $r-1 \leq a^2/(4n\sigma^2)$. Taking r to be the largest even integer less than or equal to $a^2/(6n\sigma^2)$, the tail probability is at most $e^{-r/2}$, which is at most $e \cdot e^{-a^2/(12n\sigma^2)} \leq 3 \cdot e^{-a^2/(12n\sigma^2)}$, proving the theorem.

12.6 Applications of the tail bound

Chernoff Bounds

Chernoff bounds deal with sums of Bernoulli random variables. Here we apply Theorem 12.5 to derive these.

Theorem 12.6 Suppose y_1, y_2, \ldots, y_n are independent 0-1 random variables with $E(y_i) = p$ for all *i*. Let $y = y_1 + y_2 + \cdots + y_n$. Then for any $c \in [0, 1]$,

$$Pr(|y - E(y)| \ge cnp) \le 3e^{-npc^2/8}.$$

Proof: Let $x_i = y_i - p$. Then, $E(x_i) = 0$ and $E(x_i^2) = E(y - p)^2 = p$. For $s \ge 3$,

$$|E(x_i^s)| = |E(y_i - p)^s|$$

= $|p(1 - p)^s + (1 - p)(0 - p)^s|$
= $|p(1 - p)((1 - p)^{s-1} + (-p)^{s-1})|$
 $\leq p.$

Apply Theorem 12.5 with a = cnp. Noting that $a < \sqrt{2} np$, completes the proof.

Section (12.4.11) contains a different proof that uses a standard method based on moment-generating functions and gives a better constant in the exponent.

Power Law Distributions

The power law distribution of order k where k is a positive integer is

$$f(x) = \frac{k-1}{x^k} \quad \text{for} \quad x \ge 1.$$

If a random variable x has this distribution for $k \geq 4$, then

$$\mu = E(x) = \frac{k-1}{k-2}$$
 and $\operatorname{Var}(x) = \frac{k-1}{(k-2)^2(k-3)}$.

Theorem 12.7 Suppose x_1, x_2, \ldots, x_n are *i.i.d*, each distributed according to the Power Law of order $k \ge 4$ (with $n > 10k^2$). Then, for $x = x_1 + x_2 + \cdots + x_n$, and any $\varepsilon \in (1/(2\sqrt{nk}), 1/k^2)$, we have

$$Pr(|x - E(x)| \ge \varepsilon E(x)) \le \left(\frac{4}{\varepsilon^2(k-1)n}\right)^{(k-3)/2}$$

Proof: For integer s, the s^{th} moment of $x_i - E(x_i)$, namely, $E((x_i - \mu)^s)$, exists if and only if $s \le k - 2$. For $s \le k - 2$,

$$E((x_i - \mu)^s) = (k - 1) \int_1^\infty \frac{(y - \mu)^s}{y^k} dy$$

Using the substitution of variable $z = \mu/y$

$$\frac{(y-\mu)^s}{y^k} = y^{s-k}(1-z)^s = \frac{z^{k-s}}{\mu^{k-s}}(1-z)^s$$

As y goes from 1 to ∞ , z goes from μ to 0, and $dz = -\frac{\mu}{y^2} dy$. Thus

$$\begin{split} E((x_i - \mu)^s) = & (k - 1) \int_1^\infty \frac{(y - \mu)^s}{y^k} dy \\ = & \frac{k - 1}{\mu^{k - s - 1}} \int_0^1 (1 - z)^s z^{k - s - 2} dz + \frac{k - 1}{\mu^{k - s - 1}} \int_1^\mu (1 - z)^s z^{k - s - 2} dz \end{split}$$

The first integral is just the standard integral of the beta function and its value is $\frac{s!(k-2-s)!}{(k-1)!}$. To bound the second integral, note that for $z \in [1, \mu], |z-1| \leq \frac{1}{k-2}$ and

$$z^{k-s-2} \le \left(1 + \left(1/(k-2)\right)\right)^{k-s-2} \le e^{(k-s-2)/(k-2)} \le e.$$

So, $|E((x_i - \mu)^s)| \le \frac{(k-1)s!(k-2-s)!}{(k-1)!} + \frac{e(k-1)}{(k-2)^{s+1}} \le s! \operatorname{Var}(y) \left(\frac{1}{k-4} + \frac{e}{3!}\right) \le s! \operatorname{Var}(x).$

Now, apply the first inequality of Theorem 12.5 with s of that theorem set to k-2 or k-3 whichever is even. Note that $a = \varepsilon E(x) \le \sqrt{2}n\sigma^2$ (since $\varepsilon \le 1/k^2$). The present theorem follows by a calculation.

12.7 Eigenvalues and Eigenvectors

Let A be an $n \times n$ real matrix. The scalar λ is called an eigenvalue of A if there exists a nonzero vector \mathbf{x} satisfying the equation $A\mathbf{x} = \lambda \mathbf{x}$. The vector \mathbf{x} is called the eigenvector of A associated with λ . The set of all eigenvectors associated with a given eigenvalue form a subspace as seen from the fact that if $A\mathbf{x} = \lambda \mathbf{x}$ and $A\mathbf{y} = \lambda \mathbf{y}$, then for any scalers c and d, $A(c\mathbf{x} + d\mathbf{y}) = \lambda(c\mathbf{x} + d\mathbf{y})$. The equation $A\mathbf{x} = \lambda \mathbf{x}$ has a nontrivial solution only if $\det(A - \lambda I) = 0$. The equation $\det(A - \lambda I) = 0$ is called the *characteristic* equation and has n not necessarily distinct roots.

Matrices A and B are similar if there is an invertible matrix P such that $A = P^{-1}BP$.

Theorem 12.8 If A and B are similar, then they have the same eigenvalues.

Proof: Let A and B be similar matrices. Then there exists an invertible matrix P such that $A = P^{-1}BP$. For an eigenvector \mathbf{x} of A with eigenvalue λ , $A\mathbf{x} = \lambda \mathbf{x}$, which implies $P^{-1}BP\mathbf{x} = \lambda \mathbf{x}$ or $B(P\mathbf{x}) = \lambda(P\mathbf{x})$. So, $P\mathbf{x}$ is an eigenvector of B with the same eigenvalue λ . Since the reverse also holds, the theorem follows.

Even though two similar matrices, A and B, have the same eigenvalues, their eigenvectors are in general different.

The matrix A is *diagonalizable* if A is similar to a diagonal matrix.

Theorem 12.9 A is diagonalizable if and only if A has n linearly independent eigenvectors.

Proof:

(only if) Assume A is diagonalizable. Then there exists an invertible matrix P and a diagonal matrix D such that $D = P^{-1}AP$. Thus, PD = AP. Let the diagonal elements of D be $\lambda_1, \lambda_2, \ldots, \lambda_n$ and let $\mathbf{p_1}, \mathbf{p_2}, \ldots, \mathbf{p_n}$ be the columns of P. Then $AP = [A\mathbf{p_1}, A\mathbf{p_2}, \ldots, A\mathbf{p_n}]$ and $PD = [\lambda_1\mathbf{p_1}, \lambda_2\mathbf{p_2}, \ldots, \lambda_n\mathbf{p_n}]$. Hence $A\mathbf{p_i} = \lambda_i\mathbf{p_i}$. That

is, the λ_i are the eigenvalues of A and the $\mathbf{p_i}$ are the corresponding eigenvectors. Since P is invertible, the $\mathbf{p_i}$ are linearly independent.

(if) Assume that A has n linearly independent eigenvectors $\mathbf{p_1}, \mathbf{p_2}, \ldots, \mathbf{p_n}$ with corresponding eigenvalues $\lambda_1, \lambda_2, \ldots, \lambda_n$. Then $A\mathbf{p_i} = \lambda_i \mathbf{p_i}$ and reversing the above steps

$$AP = [A\mathbf{p_1}, A\mathbf{p_2}, \dots, A\mathbf{p_n}] = [\lambda_1\mathbf{p_1}, \lambda_2\mathbf{p_2}, \dots, \lambda_n\mathbf{p_n}] = PD.$$

Thus, AP = DP. Since the \mathbf{p}_i are linearly independent, P is invertible and hence $A = P^{-1}DP$. Thus, A is diagonalizable.

It follows from the proof of the theorem that if A is diagonalizable and has eigenvalue λ with multiplicity k, then there are k linearly independent eigenvectors associated with λ .

A matrix P is orthogonal if it is invertible and $P^{-1} = P^T$. A matrix A is orthogonally diagonalizable if there exists an orthogonal matrix P such that $P^{-1}AP = D$ is diagonal. If A is orthogonally diagonalizable, then $A = PDP^T$ and AP = PD. Thus, the columns of P are the eigenvectors of A and the diagonal elements of D are the corresponding eigenvalues.

If P is an orthogonal matrix, then P^TAP and A are both representations of the same linear transformation with respect to different bases. To see this, note that if $\mathbf{e_1}, \mathbf{e_2}, \ldots, \mathbf{e_n}$ is the standard basis, then a_{ij} is the component of $A\mathbf{e_j}$ along the direction $\mathbf{e_i}$, namely, $a_{ij} = \mathbf{e_i}^T A \mathbf{e_j}$. Thus, A defines a linear transformation by specifying the image under the transformation of each basis vector. Denote by $\mathbf{p_j}$ the j^{th} column of P. It is easy to see that $(P^TAP)_{ij}$ is the component of $A\mathbf{p_j}$ along the direction $\mathbf{p_i}$, namely, $(P^TAP)_{ij} = \mathbf{p_i}^T A \mathbf{p_j}$. Since P is orthogonal, the $\mathbf{p_j}$ form a basis of the space and so P^TAP represents the same linear transformation as A, but in the basis p_1, p_2, \ldots, p_n .

Another remark is in order. Check that

$$A = PDP^{T} = \sum_{i=1}^{n} d_{ii} \mathbf{p_i} \mathbf{p_i}^{T}.$$

Compare this with the singular value decomposition where

$$A = \sum_{i=1}^{n} \sigma_i \mathbf{u}_i \mathbf{v}_i^T,$$

the only difference being that \mathbf{u}_i and \mathbf{v}_i can be different and indeed if A is not square, they will certainly be.

12.7.1 Symmetric Matrices

For an arbitrary matrix, some of the eigenvalues may be complex. However, for a symmetric matrix with real entries, all eigenvalues are real. The number of eigenvalues of a symmetric matrix, counting multiplicities, equals the dimension of the matrix. The set of eigenvectors associated with a given eigenvalue form a vector space. For a nonsymmetric matrix, the dimension of this space may be less than the multiplicity of the eigenvalue. Thus, a nonsymmetric matrix may not be diagonalizable. However, for a symmetric matrix the eigenvectors associated with a given eigenvalue form a vector space of dimension equal to the multiplicity of the eigenvalue. Thus, all symmetric matrices are diagonalizable. The above facts for symmetric matrices are summarized in the following theorem.

Theorem 12.10 (Real Spectral Theorem) Let A be a real symmetric matrix. Then

- 1. The eigenvalues, $\lambda_1, \lambda_2, \ldots, \lambda_n$, are real, as are the components of the corresponding eigenvectors, $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_n}$.
- 2. (Spectral Decomposition) A is orthogonally diagonalizable and indeed

$$A = VDV^T = \sum_{i=1}^n \lambda_i \mathbf{v_i v_i}^T,$$

where V is the matrix with columns $\mathbf{v_1}, \mathbf{v_2}, \ldots, \mathbf{v_n}, |\mathbf{v_i}| = 1$ and D is a diagonal matrix with entries $\lambda_1, \lambda_2, \ldots, \lambda_n$.

Proof: $A\mathbf{v}_{\mathbf{i}} = \lambda_i \mathbf{v}_{\mathbf{i}}$ and $\mathbf{v}_{\mathbf{i}}^c A \mathbf{v}_{\mathbf{i}} = \lambda_i \mathbf{v}_{\mathbf{i}}^c \mathbf{v}_{\mathbf{i}}$. Here the *c* superscript means conjugate transpose. Then

$$\lambda_i = \mathbf{v_i}^c A \mathbf{v_i} = (\mathbf{v_i}^c A \mathbf{v_i})^{cc} = (\mathbf{v_i}^c A^c \mathbf{v_i})^c = (\mathbf{v_i}^c A \mathbf{v_i})^c = \lambda_i^c$$

and hence λ_i is real.

Since λ_i is real, a nontrivial solution to $(A - \lambda_i I) \mathbf{x} = 0$ has real components.

Let P be a real symmetric matrix such that $P\mathbf{v_1} = \mathbf{e_1}$ where $\mathbf{e_1} = (1, 0, 0, \dots, 0)^T$ and $P^{-1} = P^T$. We will construct such a P shortly. Since $A\mathbf{v_1} = \lambda_1 \mathbf{v_1}$,

$$PAP^T \mathbf{e_1} = PAv_1 = \lambda Pv_1 = \lambda_1 \mathbf{e_1}.$$

The condition $PAP^T \mathbf{e_1} = \lambda_1 \mathbf{e_1}$ plus symmetry implies that $PAP^T = \begin{pmatrix} \lambda_1 & 0 \\ 0 & A' \end{pmatrix}$ where A' is n-1 by n-1 and symmetric. By induction, A' is orthogonally diagonalizable. Let Q be the orthogonal matrix with $QA'Q^T = D'$, a diagonal matrix. Q is $(n-1) \times (n-1)$. Augment Q to an $n \times n$ matrix by putting 1 in the (1, 1) position and 0 elsewhere in the first row and column. Call the resulting matrix R. R is orthogonal too.

$$R\left(\begin{array}{cc}\lambda_1 & 0\\ 0 & A'\end{array}\right)R^T = \left(\begin{array}{cc}\lambda_1 & 0\\ 0 & D'\end{array}\right) \implies RPAP^TR^T = \left(\begin{array}{cc}\lambda_1 & 0\\ 0 & D'\end{array}\right).$$

Since the product of two orthogonal matrices is orthogonal, this finishes the proof of (2) except it remains to construct P. For this, take an orthonormal basis of space containing $\mathbf{v_1}$. Suppose the basis is $\{\mathbf{v_1}, \mathbf{w_2}, \mathbf{w_3}, \ldots\}$ and V is the matrix with these basis vectors as its columns. Then $P = V^T$ will do.

Theorem 12.11 (The fundamental theorem of symmetric matrices) A real matrix A is orthogonally diagonalizable if and only if A is symmetric.

Proof: (if) Assume A is orthogonally diagonalizable. Then there exists P such that $D = P^{-1}AP$. Since $P^{-1} = P^T$, we get

$$A = PDP^{-1} = PDP^T$$

which implies

$$A^T = (PDP^T)^T = PDP^T = A$$

and hence A is symmetric. (only if) Already roved.

Note that a nonsymmetric matrix may not be diagonalizable, it may have eigenvalues that are not real, and the number of linearly independent eigenvectors corresponding to an eigenvalue may be less than its multiplicity. For example, the matrix

$$\left(\begin{array}{rrrr} 1 & 1 & 0 \\ 0 & 1 & 1 \\ 1 & 0 & 1 \end{array}\right)$$

has eigenvalues 2, $\frac{1}{2} + i\frac{\sqrt{3}}{2}$, and $\frac{1}{2} - i\frac{\sqrt{3}}{2}$. The matrix $\begin{pmatrix} 1 & 2 \\ 0 & 1 \end{pmatrix}$ has characteristic equation $(1 - \lambda)^2 = 0$ and thus has eigenvalue 1 with multiplicity 2 but has only one linearly independent eigenvector associated with the eigenvalue 1, namely $\mathbf{x} = c \begin{pmatrix} 1 \\ 0 \end{pmatrix}$ $c \neq 0$. Neither of these situations is possible for a symmetric matrix.

12.7.2 Relationship between SVD and Eigen Decomposition

The singular value decomposition exists for any $n \times d$ matrix whereas the eigenvalue decomposition exists only for certain square matrices. For symmetric matrices the decompositions are essentially the same.

The singular values of a matrix are always positive since they are the sum of squares of the projection of a row of a matrix onto a singular vector. Given a symmetric matrix, the eigenvalues can be positive or negative. If A is a symmetric matrix with eigenvalue decomposition $A = V_E D_E V_E^T$ and singular value decomposition $A = U_S D_S V_S^T$, what is the relationship between D_E and D_S , and between V_E and V_S , and between U_S and V_E ? Observe that if A can be expressed as QDQ^T where Q is orthonormal and D is diagonal, then AQ = QD. That is, each column of Q is an eigenvector and the elements of D are the eigenvalues. Thus, if the eigenvalues of A are distinct, then Q is unique up to a permutation of columns. If an eigenvalue has multiplicity k, then the space spanned the k columns is unique. In the following we will use the term essentially unique to

capture this situation. Now $AA^T = U_S D_S^2 U_S^T$ and $A^T A = V_S D_S^2 V_S^T$. By an argument similar to the one above, U_S and V_S are essentially unique and are the eigenvectors or negatives of the eigenvectors of A and A^T . The eigenvalues of AA^T or $A^T A$ are the squares of the eigenvalues of A. If A is not positive semi definite and has negative eigenvalues, then in the singular value decomposition $A = U_S D_S V_S$, some of the left singular vectors are the negatives of the eigenvectors. Let S be a diagonal matrix with $\pm 1's$ on the diagonal depending on whether the corresponding eigenvalue is positive or negative. Then $A = (U_S S)(SD_S)V_S$ where $U_S S = V_E$ and $SD_S = D_E$.

12.7.3 Extremal Properties of Eigenvalues

In this section we derive a min max characterization of eigenvalues that implies that the largest eigenvalue of a symmetric matrix A has a value equal to the maximum of $\mathbf{x}^T A \mathbf{x}$ over all vectors \mathbf{x} of unit length. That is, the largest eigenvalue of A equals the 2-norm of A. If A is a real symmetric matrix there exists an orthogonal matrix P that diagonalizes A. Thus

$$P^T A P = D$$

where D is a diagonal matrix with the eigenvalues of A, $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_n$, on its diagonal. Rather than working with A, it is easier to work with the diagonal matrix D. This will be an important technique that will simplify many proofs.

Consider maximizing $\mathbf{x}^T A \mathbf{x}$ subject to the conditions

 $\leq s$

1.
$$\sum_{i=1}^{n} x_i^2 = 1$$

2. $\mathbf{r}_i^T \mathbf{x} = 0, \quad 1 \le i$

where the r_i are any set of nonzero vectors. We ask over all possible sets $\{r_i | 1 \le i \le s\}$ of s vectors, what is the minimum value assumed by this maximum.

Theorem 12.12 (Min max theorem) For a symmetric matrix A, $\min_{\mathbf{r}_1,...,\mathbf{r}_s} \max_{\mathbf{r}_i \perp \mathbf{x}}^{\mathbf{x}}(\mathbf{x}^t A \mathbf{x}) =$

 λ_{s+1} where the minimum is over all sets $\{r_1, r_2, \ldots, r_s\}$ of s nonzero vectors and the maximum is over all unit vectors x orthogonal to the s nonzero vectors.

Proof: A is orthogonally diagonalizable. Let P satisfy $P^T P = I$ and $P^T A P = D$, D diagonal. Let $\mathbf{y} = P^T \mathbf{x}$. Then $\mathbf{x} = P \mathbf{y}$ and

$$\mathbf{x}^{T}A\mathbf{x} = \mathbf{y}^{T}P^{T}AP\mathbf{y} = \mathbf{y}^{T}D\mathbf{y} = \sum_{i=1}^{n} \lambda_{i}y_{i}^{2}$$

Since there is a one-to-one correspondence between unit vectors \mathbf{x} and \mathbf{y} , maximizing $\mathbf{x}^T A \mathbf{x}$ subject to $\sum x_i^2 = 1$ is equivalent to maximizing $\sum_{i=1}^n \lambda_i y_i^2$ subject to $\sum y_i^2 = 1$. Since

 $\lambda_1 \geq \lambda_i, 2 \leq i \leq n, \mathbf{y} = (1, 0, \dots, 0)$ maximizes $\sum_{i=1}^n \lambda_i y_i^2$ at λ_1 . Then $\mathbf{x} = P\mathbf{y}$ is the first column of P and is the first eigenvector of A. Similarly λ_n is the minimum value of $\mathbf{x}^T A \mathbf{x}$ subject to the same conditions.

Now consider maximizing $\mathbf{x}^T A \mathbf{x}$ subject to the conditions

1. $\sum x_i^2 = 1$ 2. $\mathbf{r}_i^T \mathbf{x} = 0$

where the \mathbf{r}_i are any set of nonzero vectors. We ask over all possible choices of s vectors what is the minimum value assumed by this maximum.

$$\min_{\mathbf{r}_1,\ldots,\mathbf{r}_s} \max_{\mathbf{r}_1^T \mathbf{x} = 0} \mathbf{x}^T A \mathbf{x}$$

As above, we may work with **y**. The conditions are

1.
$$\sum y_i^2 = 1$$

2. $\mathbf{q}_i^T \mathbf{y} = 0$ where, $\mathbf{q}_i^T = \mathbf{r}_i^T P$

Consider any choice for the vectors $\mathbf{r}_1, \mathbf{r}_2, \ldots, \mathbf{r}_s$. This gives a corresponding set of \mathbf{q}_i . The \mathbf{y}_i therefore satisfy s linear homogeneous equations. If we add $y_{s+2} = y_{s+3} = \cdots + y_n = 0$ we have n-1 homogeneous equations in n unknowns y_1, \ldots, y_n . There is at least one solution that can be normalized so that $\sum y_i^2 = 1$. With this choice of \mathbf{y}

$$\mathbf{y}^T D \mathbf{y} = \sum \lambda_i y_i^2 \ge \lambda_{s+1}$$

since coefficients greater than or equal to s + 1 are zero. Thus, for any choice of $\mathbf{r_i}$ there will be a **y** such that

$$\max_{\substack{\mathbf{y}\\\mathbf{r}_{i}^{T}\mathbf{y}=0}} \left(\mathbf{y}^{T} P^{T} A P \mathbf{y} \right) \ge \lambda_{s+1}$$

and hence

$$\min_{\mathbf{r_1},\mathbf{r_2},\ldots,\mathbf{r_s}} \max_{\substack{\mathbf{y} \\ \mathbf{r}_i^T \mathbf{y} = 0}} (\mathbf{y}^T P^T A P \mathbf{y}) \ge \lambda_{s+1}.$$

However, there is a set of s constraints for which the minimum is less than or equal to λ_{s+1} . Fix the relations to be $y_i = 0$, $1 \le i \le s$. There are s equations in n unknowns and for any y subject to these relations

$$\mathbf{y}^T D \mathbf{y} = \sum_{s+1}^n \lambda_i y_i^2 \le \lambda_{s+1}.$$

Combining the two inequalities, min max $\mathbf{y}^T D \mathbf{y} = \lambda_{s+1}$.

The above theorem tells us that the maximum of $\mathbf{x}^T A \mathbf{x}$ subject to the constraint that $|\mathbf{x}|^2 = 1$ is λ_1 . Consider the problem of maximizing $\mathbf{x}^T A \mathbf{x}$ subject to the additional restriction that \mathbf{x} is orthogonal to the first eigenvector. This is equivalent to maximizing $\mathbf{y}^t P^t A P \mathbf{y}$ subject to \mathbf{y} being orthogonal to $(1, 0, \ldots, 0)$, i.e. the first component of \mathbf{y} being 0. This maximum is clearly λ_2 and occurs for $\mathbf{y} = (0, 1, 0, \ldots, 0)$. The corresponding \mathbf{x} is the second column of P or the second eigenvector of A.

Similarly the maximum of $\mathbf{x}^T A \mathbf{x}$ for $\mathbf{p_1}^T \mathbf{x} = \mathbf{p_2}^T \mathbf{x} = \cdots \mathbf{p_s}^T \mathbf{x} = 0$ is λ_{s+1} and is obtained for $\mathbf{x} = \mathbf{p_{s+1}}$.

12.7.4 Eigenvalues of the Sum of Two Symmetric Matrices

The min max theorem is useful in proving many other results. The following theorem shows how adding a matrix B to a matrix A changes the eigenvalues of A. The theorem is useful for determining the effect of a small perturbation on the eigenvalues of A.

Theorem 12.13 Let A and B be $n \times n$ symmetric matrices. Let C=A+B. Let α_i , β_i , and γ_i denote the eigenvalues of A, B, and C respectively, where $\alpha_1 \geq \alpha_2 \geq \ldots \alpha_n$ and similarly for β_i, γ_i . Then $\alpha_s + \beta_1 \geq \gamma_s \geq \alpha_s + \beta_n$.

Proof: By the min max theorem we have

$$\alpha_s = \min_{\mathbf{r}_1, \dots, \mathbf{r}_{s-1}} \max_{\substack{\mathbf{x} \\ \mathbf{r}_i \perp \mathbf{x}}} (\mathbf{x}^T A \mathbf{x}).$$

Suppose $\mathbf{r_1}, \mathbf{r_2}, \ldots, \mathbf{r_{s-1}}$ attain the minimum in the expression. Then using the min max theorem on C,

$$\gamma_{s} \leq \max_{\mathbf{x}\perp\mathbf{r}_{1},\mathbf{r}_{2},\ldots\mathbf{r}_{s-1}} (\mathbf{x}^{T}(A+B)\mathbf{x})$$

$$\leq \max_{\mathbf{x}\perp\mathbf{r}_{1},\mathbf{r}_{2},\ldots\mathbf{r}_{s-1}} (\mathbf{x}^{T}A\mathbf{x}) + \max_{\mathbf{x}\perp\mathbf{r}_{1},\mathbf{r}_{2},\ldots\mathbf{r}_{s-1}} (\mathbf{x}^{T}B\mathbf{x})$$

$$\leq \alpha_{s} + \max_{\mathbf{x}} (\mathbf{x}^{T}B\mathbf{x}) \leq \alpha_{s} + \beta_{1}.$$

Therefore, $\gamma_s \leq \alpha_s + \beta_1$.

An application of the result to A = C + (-B), gives $\alpha_s \leq \gamma_s - \beta_n$. The eigenvalues of -B are minus the eigenvalues of B and thus $-\beta_n$ is the largest eigenvalue. Hence $\gamma_s \geq \alpha_s + \beta_n$ and combining inequalities yields $\alpha_s + \beta_1 \geq \gamma_s \geq \alpha_s + \beta_n$.

Lemma 12.14 Let A and B be $n \times n$ symmetric matrices. Let C=A+B. Let α_i , β_i , and γ_i denote the eigenvalues of A, B, and C respectively, where $\alpha_1 \geq \alpha_2 \geq \ldots \alpha_n$ and similarly for β_i, γ_i . Then $\gamma_{r+s-1} \leq \alpha_r + \beta_s$.

Proof: There is a set of r-1 relations such that over all **x** satisfying the r-1 relationships

$$\max(\mathbf{x}^T A \mathbf{x}) = \alpha_r.$$

And a set of s-1 relations such that over all **x** satisfying the s-1 relationships

$$\max(\mathbf{x}^T B \mathbf{x}) = \beta_s.$$

Consider **x** satisfying all these r + s - 2 relations. For any such **x**

$$\mathbf{x}^T C \mathbf{x} = \mathbf{x}^T A \mathbf{x} + \mathbf{x}^T B \mathbf{x} \mathbf{x} \le \alpha_r + \beta_s$$

and hence over all the ${\bf x}$

$$\max(\mathbf{x}^T C \mathbf{x}) \le \alpha_s + \beta_r$$

Taking the minimum over all sets of r + s - 2 relations

$$\gamma_{r+s-1} = \min\max(\mathbf{x}^T C \mathbf{x}) \le \alpha_r + \beta_s$$

A set of vectors $\{\mathbf{x_1}, \ldots, \mathbf{x_n}\}$ is orthogonal if $\mathbf{x_i}^T \mathbf{x_j} = 0$ for $i \neq j$ and is orthonormal if in addition $|\mathbf{x_i}| = 1$ for all *i*. A matrix *A* is orthonormal if $A^T A = I$. If *A* is a square orthonormal matrix, then rows as well as columns are orthogonal. In other words, if *A* is square orthonormal, then A^T is also. In the case of matrices over the complexes, the concept of an orthonormal matrix is replaced by that of a unitary matrix. A^* is the conjugate transpose of *A* if $a_{ij}^* = \bar{a}_{ji}$ where a_{ij}^* is the ij^{th} entry of A^* and \bar{a}_{ij}^* is the complex conjugate of the ij^{th} element of *A*. A matrix *A* over the field of complex numbers is **unitary** if $AA^* = I$.

Norms

A norm on \mathbb{R}^n is a function $f: \mathbb{R}^n \to \mathbb{R}$ satisfying the following three axioms:

- 1. $f(\mathbf{x}) \ge 0$,
- 2. $f(\mathbf{x} + \mathbf{y}) \leq f(\mathbf{x}) + f(\mathbf{y})$, and
- 3. $f(\alpha \mathbf{x}) = |\alpha| f(\mathbf{x}).$

A norm on a vector space provides a distance function where

distance
$$(\mathbf{x}, \mathbf{y}) = norm(\mathbf{x} - \mathbf{y}).$$

An important class of norms for vectors is the *p*-norms defined for p > 0 by

$$|\mathbf{x}|_p = (|\mathbf{x}_1|^p + \dots + |\mathbf{x}_n|^p)^{\frac{1}{p}}$$

Important special cases are

 $|\mathbf{x}|_0$ the number of non zero entries $|\mathbf{x}|_1 = |x_1| + \dots + |x_n|$

$$\begin{aligned} \mathbf{x}|_{1} &= |\mathbf{x}_{1}|^{2} + \cdots + |\mathbf{x}_{n}|^{2} \\ \mathbf{x}|_{2} &= \sqrt{|x_{1}|^{2} + \cdots + |x_{n}|^{2}} \\ \mathbf{x}|_{\infty} &= \max |x_{i}|. \end{aligned}$$

Lemma 12.15 For any $1 \le p < q$, $|\mathbf{x}|_q \le |\mathbf{x}|_p$.

Proof:

$$|\mathbf{x}|_q^q = \sum_i |x_i|^q.$$

Let $a_i = |x_i|^q$ and $\rho = p/q$. Using Jensen's inequality (see Section 12.3) that for any nonnegative reals a_1, a_2, \ldots, a_n and any $\rho \in (0, 1)$, we have $(\sum_{i=1}^n a_i)^{\rho} \leq \sum_{i=1}^n a_i^{\rho}$, the lemma is proved.

There are two important matrix norms, the matrix p-norm

$$\left|\left|A\right|\right|_{p} = \max_{|\mathbf{x}|=1} \left\|A\mathbf{x}\right\|_{p}$$

and the Frobenius norm

$$||A||_F = \sqrt{\sum_{ij} a_{ij}^2}.$$

Let $\mathbf{a}_{\mathbf{i}}$ be the i^{th} column of A. Then $||A||_F^2 = \sum_i \mathbf{a}_i^T \mathbf{a}_i = tr(A^T A)$. A similar argument on the rows yields $||A||_F^2 = tr(AA^T)$. Thus, $||A||_F^2 = tr(A^T A) = tr(AA^T)$. If A is symmetric and rank k

$$||A||_{2}^{2} \leq ||A||_{F}^{2} \leq k ||A||_{2}^{2}.$$

12.7.6 Important Norms and Their Properties

Lemma 12.16 $||AB||_2 \le ||A||_2 ||B||_2$

Proof: $||AB||_2 = \max_{|\mathbf{x}|=1} |AB\mathbf{x}|$. Let \mathbf{y} be the value of \mathbf{x} that achieves the maximum and let $\mathbf{z} = B\mathbf{y}$. Then

$$||AB||_2 = |AB\mathbf{y}| = |A\mathbf{z}| = \left|A\frac{\mathbf{z}}{|\mathbf{z}|}\right||\mathbf{z}|$$

But $\left|A_{\frac{\mathbf{z}}{|\mathbf{z}|}}\right| \le \max_{|\mathbf{x}|=1} |A\mathbf{x}| = ||A||_2$ and $|\mathbf{z}| \le \max_{|\mathbf{x}|=1} |B\mathbf{x}| = ||B||_2$. Thus $||AB||_2 \le ||A||_2 ||B||_2$.

Let Q be an orthonormal matrix.

Lemma 12.17 For all \mathbf{x} , $|Q\mathbf{x}| = |\mathbf{x}|$. Proof: $|Q\mathbf{x}|_2^2 = \mathbf{x}^T Q^T Q \mathbf{x} = \mathbf{x}^T \mathbf{x} = |\mathbf{x}|_2^2$.

Lemma 12.18 $||QA||_2 = ||A||_2$

Proof: For all \mathbf{x} , $|Q\mathbf{x}| = |\mathbf{x}|$. Replacing \mathbf{x} by $A\mathbf{x}$, $|QA\mathbf{x}| = |A\mathbf{x}|$ and thus $\max_{|\mathbf{x}|=1} |QA\mathbf{x}| = \max_{|\mathbf{x}|=1} |A\mathbf{x}|$

Lemma 12.19 $||AB||_F^2 \le ||A||_F^2 ||B||_F^2$

Proof: Let \mathbf{a}_i be the i^{th} column of A and let \mathbf{b}_j be the j^{th} column of B. By the Cauchy-Schwartz inequality $\|\mathbf{a}_i^T \mathbf{b}_j\| \leq \|\mathbf{a}_i\| \|\mathbf{b}_j\|$. Thus $\||AB||_F^2 = \sum_i \sum_j |\mathbf{a}_i^T \mathbf{b}_j|^2 \leq$

$$\sum_{i} \sum_{j} \|\mathbf{a}_{i}\|^{2} \|\mathbf{b}_{j}\|^{2} = \sum_{i} \|\mathbf{a}_{i}\|^{2} \sum_{j} \|\mathbf{b}_{j}\|^{2} = \|A\|_{F}^{2} \|B\|$$

Lemma 12.20 $||QA||_F = ||A||_F$

Proof: $||QA||_F^2 = \text{Tr}(A^T Q^T Q A) = \text{Tr}(A^T A) = ||A||_F^2.$

Lemma 12.21 For real, symmetric matrix A with eigenvalues $\lambda_1 \geq \lambda_2 \geq \ldots$, $||A||_2^2 = \max(\lambda_1^2, \lambda_n^2)$ and $||A||_F^2 = \lambda_1^2 + \lambda_2^2 + \cdots + \lambda_n^2$

Proof: Suppose the spectral decomposition of A is PDP^T , where P is an orthogonal matrix and D is diagonal. We saw that $||P^TA||_2 = ||A||_2$. Applying this again, $||P^TAP||_2 = ||A||_2$. But, $P^TAP = D$ and clearly for a diagonal matrix D, $||D||_2$ is the largest absolute value diagonal entry from which the first equation follows. The proof of the second is analogous.

If A is real and symmetric and of rank k then $||A||_2^2 \leq ||A||_F^2 \leq k ||A||_2^2$

Theorem 12.22 $||A||_2^2 \le ||A||_F^2 \le k ||A||_2^2$

Proof: It is obvious for diagonal matrices that $||D||_2^2 \leq ||D||_F^2 \leq k ||D||_2^2$. Let $D = Q^t A Q$ where Q is orthonormal. The result follows immediately since for Q orthonormal, $||QA||_2 = ||A||_2$ and $||QA||_F = ||A||_F$.

Real and symmetric are necessary for some of these theorems. This condition was needed to express $\Sigma = Q^T A Q$. For example, in Theorem 12.22 suppose A is the $n \times n$ matrix

$$A = \left(\begin{array}{rrrr} 1 & 1 & & \\ 1 & 1 & & \\ \vdots & \vdots & 0 \\ 1 & 1 & & \end{array} \right).$$

 $||A||_2 = 2$ and $||A||_F = \sqrt{2n}$. But A is rank 2 and $||A||_F > 2 ||A||_2$ for n > 8.

Lemma 12.23 Let A be a symmetric matrix. Then $||A||_2 = \max_{|\mathbf{x}|=1} |\mathbf{x}^T A \mathbf{x}|$.

Proof: By definition, the 2-norm of A is $||A||_2 = \max_{|\mathbf{x}|=1} |A\mathbf{x}|$. Thus,

$$\|A\|_{2} = \max_{|\mathbf{x}|=1} |A\mathbf{x}| = \max_{|\mathbf{x}|=1} \sqrt{\mathbf{x}^{T} A^{T} A \mathbf{x}} = \sqrt{\lambda_{1}^{2}} = \lambda_{1} = \max_{|\mathbf{x}|=1} |\mathbf{x}^{T} A \mathbf{x}|$$

The two norm of a matrix A is greater than or equal to the 2-norm of any of its columns. Let $\mathbf{a}_{\mathbf{u}}$ be a column of A.

Lemma 12.24 $|\mathbf{a}_{\mathbf{u}}| \le ||A||_2$

Proof: Let $\mathbf{e}_{\mathbf{u}}$ be the unit vector with a 1 in position u and all other entries zero. Note $\lambda = \max_{|x|=1} |Ax|$. Let $\mathbf{x} = \mathbf{e}_{\mathbf{u}}$ where $\mathbf{a}_{\mathbf{u}}$ is row u. Then $|\mathbf{a}_{\mathbf{u}}| = |A\mathbf{e}_{\mathbf{u}}| \le \max_{|\mathbf{x}|=1} |A\mathbf{x}| = \lambda$

12.7.7 Linear Algebra

Lemma 12.25 Let A be an $n \times n$ symmetric matrix. Then $det(A) = \lambda_1 \lambda_2 \cdots \lambda_n$.

Proof: The det $(A - \lambda I)$ is a polynomial in λ of degree n. The coefficient of λ^n will be ± 1 depending on whether n is odd or even. Let the roots of this polynomial be $\lambda_1, \lambda_2, \ldots, \lambda_n$. Then det $(A - \lambda I) = (-1)^n \prod_{i=1}^n (\lambda - \lambda_i)$. Thus

$$\det(A) = \det(A - \lambda I)|_{\lambda=0} = (-1)^n \prod_{i=1}^n (\lambda - \lambda_i) \bigg|_{\lambda=0} = \lambda_1 \lambda_2 \cdots \lambda_n$$

The trace of a matrix is defined to be the sum of its diagonal elements. That is, tr $(A) = a_{11} + a_{22} + \cdots + a_{nn}$.

Lemma 12.26 $tr(A) = \lambda_1 + \lambda_2 + \cdots + \lambda_n$.

Proof: Consider the coefficient of λ^{n-1} in $\det(A - \lambda I) = (-1)^n \prod_{i=1}^n (\lambda - \lambda_i)$. Write

$$A - \lambda I = \begin{pmatrix} a_{11} - \lambda & a_{12} & \cdots \\ a_{21} & a_{22} - \lambda & \cdots \\ \vdots & \vdots & \vdots \end{pmatrix}.$$

Calculate $\det(A - \lambda I)$ by expanding along the first row. Each term in the expansion involves a determinant of size n - 1 which is a polynomial in λ of deg n - 2 except for the principal minor which is of deg n - 1. Thus the term of deg n - 1 comes from

$$(a_{11} - \lambda) (a_{22} - \lambda) \cdots (a_{nn} - \lambda)$$

and has coefficient $(-1)^{n-1} (a_{11} + a_{22} + \dots + a_{nn})$. Now

$$(-1)^{n} \prod_{i=1}^{n} (\lambda - \lambda_{i}) = (-1)^{n} (\lambda - \lambda_{1})(\lambda - \lambda_{2}) \cdots (\lambda - \lambda_{n})$$
$$= (-1)^{n} \left(\lambda^{n} - (\lambda_{1} + \lambda_{2} + \dots + \lambda_{n})\lambda^{n-1} + \dots + \lambda_{n}\right)$$

Therefore equating coefficients $\lambda_1 + \lambda_2 + \dots + \lambda_n = a_{11} + a_{22} + \dots + a_{nn} = tr(A)$

Note that $(tr(A))^2 \neq tr(A^2)$. For example $A = \begin{pmatrix} 1 & 0 \\ 0 & 2 \end{pmatrix}$ has trace 3, $A^2 = \begin{pmatrix} 1 & 0 \\ 0 & 4 \end{pmatrix}$ has trace $5 \neq 9$. However $tr(A^2) = \lambda_1^2 + \lambda_2^2 + \dots + \lambda_n^2$. To see this, observe that $A^2 = (V^T D V)^2 = V^T D^2 V$. Thus, the eigenvalues of A^2 are the squares of the eigenvalues for A.

Alternative proof that $tr(A) = \lambda_1 + \lambda_2 + \cdots + \lambda_n$. Suppose the spectral decomposition of A is $A = PDP^T$. We have

$$\operatorname{tr}(A) = \operatorname{tr}(PDP^{T}) = \operatorname{tr}(DP^{T}P) = \operatorname{tr}(D) = \lambda_{1} + \lambda_{2} + \dots + \lambda_{n}.$$

Lemma 12.27 If A is $n \times m$ and B is a $m \times n$ matrix, then tr(AB) = tr(BA).

$$\operatorname{tr}(AB) = \sum_{i=1}^{n} \sum_{j=1}^{n} a_{ij} b_{ji} = \sum_{j=1}^{n} \sum_{i=1}^{n} b_{ji} a_{ij} = \operatorname{tr}(BA)$$

Pseudo inverse

Let A be an $n \times m$ rank r matrix and let $A = U\Sigma V^T$ be the singular value decomposition of A. Let $\Sigma' = diag\left(\frac{1}{\sigma_1}, \ldots, \frac{1}{\sigma_r}, 0, \ldots, 0\right)$ where $\sigma_1, \ldots, \sigma_r$ are the nonzero singular values of A. Then $A' = V\Sigma'U^T$ is the pseudo inverse of A. It is the unique X that minimizes $||AX - I||_F$. Second eigenvector

Suppose the eigenvalues of a matrix are $\lambda_1 \geq \lambda_2 \geq \cdots$. The second eigenvalue, λ_2 , plays an important role for matrices representing graphs. It may be the case that $|\lambda_n| > |\lambda_2|$.

Why is the second eigenvalue so important? Consider partitioning the vertices of a regular degree d graph G = (V, E) into two blocks of equal size so as to minimize the number of edges between the two blocks. Assign value +1 to the vertices in one block and -1 to the vertices in the other block. Let \mathbf{x} be the vector whose components are the ± 1 values assigned to the vertices. If two vertices, i and j, are in the same block, then x_i and x_j are both +1 or both -1 and $(x_i - x_j)^2 = 0$. If vertices i and j are in different blocks then $(x_i - x_j)^2 = 4$. Thus, partitioning the vertices into two blocks so as to minimize the edges

between vertices in different blocks is equivalent to finding a vector \mathbf{x} with coordinates ± 1 of which half of its coordinates are +1 and half of which are -1 that minimizes

$$E_{cut} = \frac{1}{4} \sum_{(i,j) \in E} (x_i - x_j)^2$$

Let A be the adjacency matrix of G. Then

$$\mathbf{x}^{T} A \mathbf{x} = \sum_{ij} a_{ij} x_{i} x_{j} = 2 \sum_{edges} x_{i} x_{j}$$

= 2 × $\begin{pmatrix} \text{number of edges} \\ \text{within components} \end{pmatrix} - 2 × \begin{pmatrix} \text{number of edges} \\ \text{between components} \end{pmatrix}$
= 2 × $\begin{pmatrix} \text{total number} \\ \text{of edges} \end{pmatrix} - 4 × \begin{pmatrix} \text{number of edges} \\ \text{between components} \end{pmatrix}$

Maximizing $\mathbf{x}^T A \mathbf{x}$ over all \mathbf{x} whose coordinates are ± 1 and half of whose coordinates are +1 is equivalent to minimizing the number of edges between components.

Since finding such an **x** is computational difficult, replace the integer condition on the components of **x** and the condition that half of the components are positive and half of the components are negative with the conditions $\sum_{i=1}^{n} x_i^2 = 1$ and $\sum_{i=1}^{n} x_i = 0$. Then finding the optimal **x** gives us the second eigenvalue since it is easy to see that the first eigenvector Is along **1**

$$\lambda_2 = \max_{\mathbf{x} \perp \mathbf{v}_1} \frac{\mathbf{x}^T A \mathbf{x}}{\sum x_i^2}$$

Actually we should use $\sum_{i=1}^{n} x_i^2 = n$ not $\sum_{i=1}^{n} x_i^2 = 1$. Thus $n\lambda_2$ must be greater than $2 \times \begin{pmatrix} \text{total number} \\ \text{of edges} \end{pmatrix} - 4 \times \begin{pmatrix} \text{number of edges} \\ \text{between components} \end{pmatrix}$ since the maximum is taken over a larger set of \mathbf{x} . The fact that λ_2 gives us a bound on the minimum number of cross edges is what makes it so important.

12.7.8 Distance between subspaces

Suppose S_1 and S_2 are two subspaces. Choose a basis of S_1 and arrange the basis vectors as the columns of a matrix X_1 ; similarly choose a basis of S_2 and arrange the basis vectors as the columns of a matrix X_2 . Note that S_1 and S_2 can have different dimensions. Define the square of the distance between two subspaces by

$$dist^{2}(S_{1}, S_{2}) = dist^{2}(X_{1}, X_{2}) = ||X_{1} - X_{2}X_{2}^{T}X_{1}||_{F}^{2}$$

Since $X_1 - X_2 X_2^T X_1$ and $X_2 X_2^T X_1$ are orthogonal

$$\|X_1\|_F^2 = \|X_1 - X_2 X_2^T X_1\|_F^2 + \|X_2 X_2^T X_1\|_F^2$$

and hence

$$dist^{2}(X_{1}, X_{2}) = \|X_{1}\|_{F}^{2} - \|X_{2}X_{2}^{T}X_{1}\|_{F}^{2}.$$

Intuitively, the distance between X_1 and X_2 is the Frobenius norm of the component of X_1 not in the space spanned by the columns of X_2 .

If X_1 and X_2 are 1-dimensional unit length vectors, $dist^2(X_1, X_2)$ is the sin squared of the angle between the spaces.

Example: Consider two subspaces in four dimensions

$$X_1 = \begin{pmatrix} \frac{1}{\sqrt{2}} & 0\\ 0 & \frac{1}{\sqrt{3}}\\ \frac{1}{\sqrt{2}} & \frac{1}{\sqrt{3}}\\ 0 & \frac{1}{\sqrt{3}} \end{pmatrix} \qquad X_2 = \begin{pmatrix} 1 & 0\\ 0 & 1\\ 0 & 0\\ 0 & 0 \end{pmatrix}$$

Here

$$dist^{2}(X_{1}, X_{2}) = \left\| \begin{pmatrix} \frac{1}{\sqrt{2}} & 0\\ 0 & \frac{1}{\sqrt{3}}\\ \frac{1}{\sqrt{2}} & \frac{1}{\sqrt{3}}\\ 0 & \frac{1}{\sqrt{3}} \end{pmatrix} - \begin{pmatrix} 1 & 0\\ 0 & 1\\ 0 & 0\\ 0 & 0 \end{pmatrix} \begin{pmatrix} 1 & 0 & 0 & 0\\ 0 & 1 & 0 & 0 \end{pmatrix} \begin{pmatrix} \frac{1}{\sqrt{2}} & 0\\ 0 & \frac{1}{\sqrt{3}}\\ \frac{1}{\sqrt{2}} & \frac{1}{\sqrt{3}}\\ 0 & \frac{1}{\sqrt{3}} \end{pmatrix} \right\|_{F}^{2}$$
$$= \left\| \begin{pmatrix} 0 & 0\\ 0 & 0\\ \frac{1}{\sqrt{2}} & \frac{1}{\sqrt{3}}\\ 0 & \frac{1}{\sqrt{3}} \end{pmatrix} \right\|_{F}^{2} = \frac{7}{6}$$

In essence, we projected each column vector of X_1 onto X_2 and computed the Frobenius norm of X_1 minus the projection. The Frobenius norm of each column is the sin squared of the angle between the original column of X_1 and the space spanned by the columns of X_2 .

12.8 Generating Functions

A sequence a_0, a_1, \ldots , can be represented by a generating function $g(x) = \sum_{i=0}^{\infty} a_i x^i$. The advantage of the generating function is that it captures the entire sequence in a closed form that can be manipulated as an entity. For example, if g(x) is the generating function for the sequence a_0, a_1, \ldots , then $x \frac{d}{dx} g(x)$ is the generating function for the sequence 0, $a_1, 2a_2, 3a_3, \ldots$ and $x^2g''(x) + xg'(x)$ is the generating function for the sequence for 0, $a_1, 4a_2, 9a_3, \ldots$

Example: The generating function for the sequence $1, 1, \ldots$ is $\sum_{i=0}^{\infty} x^i = \frac{1}{1-x}$. The generating function for the sequence $0, 1, 2, 3, \ldots$ is

$$\sum_{i=0}^{\infty} ix^{i} = \sum_{i=0}^{\infty} x \frac{d}{dx} x^{i} = x \frac{d}{dx} \sum_{i=0}^{\infty} x^{i} = x \frac{d}{dx} \frac{1}{1-x} = \frac{x}{(1-x)^{2}}.$$

Example: If A can be selected 0 or 1 times and B can be selected 0, 1, or 2 times and C can be selected 0, 1, 2, or 3 times, in how many ways can five objects be selected. Consider the generating function for the number of ways to select objects. The generating function for the number of ways of selecting objects, selecting only A's is 1+x, only B's is $1+x+x^2$, and only C's is $1+x+x^2+x^3$. The generating function when selecting A's, B's, and C's is the product.

$$(1+x)(1+x+x^2)(1+x+x^2+x^3) = 1 + 3x + 5x^2 + 6x^3 + 5x^4 + 3x^5 + x^6$$

The coefficient of x^5 is 3 and hence we can select five objects in three ways: ABBCC, ABCCC, or BBCCC.

The generating functions for the sum of random variables

Let $f(x) = \sum_{i=0}^{\infty} p_i x^i$ be the generating function for an integer valued random variable

where p_i is the probability that the random variable takes on value *i*. Let $g(x) = \sum_{i=0}^{\infty} q_i x^i$

be the generating function of an independent integer valued random variable where q_i is the probability that the random variable takes on the value *i*. The sum of these two random variables has the generating function f(x)g(x). This is because the coefficient of x^i in the product f(x)g(x) is $\sum_{k=0}^{i} p_k q_{k-i}$ and this is also the probability that the sum of the random variables is *i*. Repeating this, the generating function of a sum of independent nonnegative integer valued random variables is the product of their generating functions.

12.8.1 Generating Functions for Sequences Defined by Recurrence Relationships

Consider the Fibonacci sequence

$$0, 1, 1, 2, 3, 5, 8, 13, 21, 34, 55, 89, \ldots$$

defined by the recurrence relationship

$$f_0 = 0$$
 $f_1 = 1$ $f_i = f_{i-1} + f_{i-2}$ $i \ge 2$

Multiply each side of the recurrence by x^i and sum from *i* equals two to infinity.

$$\sum_{i=2}^{\infty} f_i x^i = \sum_{i=2}^{\infty} f_{i-1} x^i + \sum_{i=2}^{\infty} f_{i-2} x^i$$
$$f_2 x^2 + f_3 x^3 + \dots = f_1 x^2 + f_2 x^3 + \dots + f_0 x^2 + f_1 x^3 + \dots$$
$$= x \left(f_1 x + f_2 x^2 + \dots \right) + x^2 \left(f_0 + f_1 x + \dots \right)$$
(12.1)

Let

$$f(x) = \sum_{i=0}^{\infty} f_i x^i.$$
 (12.2)

Substituting (12.2) into (12.1) yields

$$f(x) - f_0 - f_1 x = x (f(x) - f_0) + x^2 f(x)$$

$$f(x) - x = x f(x) + x^2 f(x)$$

$$f(x)(1 - x - x^2) = x$$

Thus, $f(x) = \frac{x}{1-x-x^2}$ is the generating function for the Fibonacci sequence.

Note that generating functions are formal manipulations and do not necessarily converge outside some region of convergence. Consider the generating function $f(x) = \sum_{i=0}^{\infty} f_i x^i = \frac{x}{1-x-x^2}$ for the Fibonacci sequence. Using $\sum_{i=0}^{\infty} f_i x^i$,

$$f(1) = f_0 + f_1 + f_2 + \dots = \infty$$

and using $f(x) = \frac{x}{1-x-x^2}$

$$f(1) = \frac{1}{1 - 1 - 1} = -1.$$

Asymptotic behavior

To determine the asymptotic behavior of the Fibonacci sequence write

$$f(x) = \frac{x}{1 - x - x^2} = \frac{\frac{\sqrt{5}}{5}}{1 - \phi_1 x} + \frac{-\frac{\sqrt{5}}{5}}{1 - \phi_2 x}$$

where $\phi_1 = \frac{1+\sqrt{5}}{2}$ and $\phi_1 = \frac{1-\sqrt{5}}{2}$ are the reciprocals of the two roots of the quadratic $1-x-x^2 = 0$.

Then

$$f(x) = \frac{\sqrt{5}}{5} \left(1 + \phi_1 x + (\phi_1 x)^2 + \dots - \left(1 + \phi_2 x + (\phi_2 x)^2 + \dots \right) \right).$$

Thus,

$$f_n = \frac{\sqrt{5}}{5} \left(\phi_1^n - \phi_2^n \right).$$

Since $\phi_2 < 1$ and $\phi_1 > 1$, for large n, $f_n \cong \frac{\sqrt{5}}{5}\phi_1^n$. In fact, since $f_n = \frac{\sqrt{5}}{5}(\phi_1^n - \phi_2^n)$ is an integer and $\phi_2 < 1$, it must be the case that $f_n = \left\lfloor f_n + \frac{\sqrt{5}}{2}\phi_2^n \right\rfloor$. Hence $f_n = \left\lfloor \frac{\sqrt{5}}{5}\phi_1^n \right\rfloor$ for all n.

Means and standard deviations of sequences

Generating functions are useful for calculating the mean and standard deviation of a sequence. Let z be an integral valued random variable where p_i is the probability that z equals i. The expected value of z is given by $m = \sum_{i=0}^{\infty} ip_i$. Let $p(x) = \sum_{i=0}^{\infty} p_i x^i$ be the generating function for the sequence p_1, p_2, \ldots The generating function for the sequence $p_1, 2p_2, 3p_3, \ldots$ is

$$x\frac{d}{dx}p(x) = \sum_{i=0}^{\infty} ip_i x^i.$$

Thus, the expected value of the random variable z is $m = xp'(x)|_{x=1} = p'(1)$. If p was not a probability function, its average value would be $\frac{p'(1)}{p(1)}$ since we would need to normalize the area under p to one.

The second moment of z, is $E(z^2) - E^2(z)$ and can be obtained as follows.

$$\begin{aligned} x^{2} \frac{d}{dx} p(x) \Big|_{x=1} &= \sum_{i=0}^{\infty} i(i-1)x^{i} p(x) \Big|_{x=1} \\ &= \sum_{i=0}^{\infty} i^{2} x^{i} p(x) \Big|_{x=1} - \sum_{i=0}^{\infty} ix^{i} p(x) \Big|_{x=1} \\ &= E(z^{2}) - E(z). \end{aligned}$$

Thus, $\sigma^2 = E(z^2) - E^2(z) = E(z^2) - E(z) + E(z) - E^2(z) = p''(1) + p'(1) - (p'(1))^2$.

12.8.2 The Exponential Generating Function and the Moment Generating Function

Besides the ordinary generating function there are a number of other types of generating functions. One of these is the exponential generating function. Given a sequence a_0, a_1, \ldots , the associated exponential generating function is $g(x) = \sum_{i=0}^{\infty} a_i \frac{x^i}{i!}$.

Moment generating functions

The k^{th} moment of a random variable x around the point b is given by $E((x-b)^k)$. Usually the word moment is used to denote the moment around the value 0 or around the mean. In the following, we use moment to mean the moment about the origin.

The moment generating function of a random variable x is defined by

$$\Psi(t) = E(e^{tx}) = \int_{-\infty}^{\infty} e^{tx} p(x) dx$$

Replacing e^{tx} by its power series expansion $1 + tx + \frac{(tx)^2}{2!} \cdots$ gives

$$\Psi(t) = \int_{-\infty}^{\infty} \left(1 + tx + \frac{\left(tx\right)^2}{2!} + \cdots \right) p(x) dx$$

Thus, the k^{th} moment of x about the origin is k! times the coefficient of t^k in the power series expansion of the moment generating function. Hence, the moment generating function is the exponential generating function for the sequence of moments about the origin.

The moment generating function transforms the probability distribution p(x) into a function $\Psi(t)$ of t. Note $\Psi(0) = 1$ and is the area or integral of p(x). The moment generating function is closely related to the *characteristic function* which is obtained by replacing e^{tx} by e^{itx} in the above integral where $i = \sqrt{-1}$ and is related to the *Fourier transform* which is obtained by replacing e^{tx} by e^{-itx} .

 $\Psi(t)$ is closely related to the Fourier transform and its properties are essentially the same. In particular, p(x) can be uniquely recovered by an inverse transform from $\Psi(t)$. More specifically, if all the moments m_i are finite and the sum $\sum_{i=0}^{\infty} \frac{m_i}{i!} t^i$ converges absolutely in a region around the origin, then p(x) is uniquely determined.

The Gaussian probability distribution with zero mean and unit variance is given by $p(x) = \frac{1}{\sqrt{2\pi}}e^{-\frac{x^2}{2}}$. Its moments are given by

$$u_n = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} x^n e^{-\frac{x^2}{2}} dx$$
$$= \begin{cases} \frac{n!}{2^{\frac{n}{2}} \binom{n}{2}!} & \text{n even} \\ 0 & \text{n odd} \end{cases}$$

To derive the above, use integration by parts to get $u_n = (n-1)u_{n-2}$ and combine this with $u_0 = 1$ and $u_1 = 0$. The steps are as follows. Let $u = e^{-\frac{x^2}{2}}$ and $v = x^{n-1}$. Then $u' = -xe^{-\frac{x^2}{2}}$ and $v' = (n-1)x^{n-2}$. Now $uv = \int u'v + \int uv'$ or

$$e^{-\frac{x^2}{2}}x^{n-1} = \int x^n e^{-\frac{x^2}{2}} dx + \int (n-1) x^{n-2} e^{-\frac{x^2}{2}} dx.$$

From which

$$\int_{-\infty}^{\infty} x^n e^{-\frac{x^2}{2}} dx = (n-1) \int_{-\infty}^{\infty} x^{n-2} e^{-\frac{x^2}{2}} dx - e^{-\frac{x^2}{2}} x^{n-1}$$
$$\int_{-\infty}^{\infty} x^n e^{-\frac{x^2}{2}} dx = (n-1) \int_{-\infty}^{\infty} x^{n-2} e^{-\frac{x^2}{2}} dx$$

Thus, $u_n = (n-1) u_{n-2}$.

The moment generating function is given by

$$g(s) = \sum_{n=0}^{\infty} \frac{u_n s^n}{n!} = \sum_{\substack{n=0\\n \text{ even}}}^{\infty} \frac{n!}{2^{\frac{n}{2}} \frac{n!}{2!}} \frac{s^n}{n!} = \sum_{i=0}^{\infty} \frac{s^{2i}}{2^i i!} = \sum_{i=0}^{\infty} \frac{1}{i!} \left(\frac{s^2}{2}\right)^i = e^{\frac{s^2}{2}}.$$

For the general Gaussian, the moment generating function is

$$g\left(s\right) = e^{su + \left(\frac{\sigma^2}{2}\right)s^2}$$

Thus, given two independent Gaussians with mean u_1 and u_2 and variances σ_1^2 and σ_2^2 , the product of their moment generating functions is

$$e^{s(u_1+u_2)+(\sigma_1^2+\sigma_2^2)s^2}$$

the moment generating function for a Gaussian with mean $u_1 + u_2$ and variance $\sigma_1^2 + \sigma_2^2$. Thus, the convolution of two Gaussians is a Gaussian and the sum of two random variables that are both Gaussian is a Gaussian random variable.

12.9 Miscellaneous

12.9.1 Lagrange multipliers

Lagrange multipliers are used to convert a constrained optimization problem into an unconstrained optimization. Suppose we wished to maximize a function $f(\mathbf{x})$ subject to a constraint $g(\mathbf{x}) = c$. The value of $f(\mathbf{x})$ along the constraint $g(\mathbf{x}) = c$ might increase for a while and then start to decrease. At the point where $f(\mathbf{x})$ stops increasing and starts to decrease, the contour line for $f(\mathbf{x})$ is tangent to the curve of the constraint $g(\mathbf{x}) = c$. Stated another way the gradient of $f(\mathbf{x})$ and the gradient of $g(\mathbf{x})$ are parallel.

By introducing a new variable λ we can express the condition by $\nabla_{\mathbf{x}} f = \lambda \nabla_{\mathbf{x}} g$ and g = c. These two conditions hold if and only if

$$\nabla_{\mathbf{x}\lambda} \left(f(\mathbf{x}) + \lambda \left(g(\mathbf{x}) - c \right) \right) = 0$$

The partial with respect to λ establishes that $g(\mathbf{x}) = c$. We have converted the constrained optimization problem in x to an unconstrained problem with variables \mathbf{x} and λ .

12.9.2 Finite Fields

For a prime p and integer n there is a unique finite field with p^n elements. In Section 4.6 we used the field $GF(2^n)$, which consists of polynomials of degree less than n with coefficients over the field GF(2). In $GF(2^8)$

$$(x^7 + x^5 + x) + (x^6 + x^5 + x^4) = x^7 + x^6 = x^4 = x$$

Multiplication is modulo an irreducible polynomial. Thus

$$(x^{7} + x^{5} + x)(x^{6} + x^{5} + x^{4}) = x^{13} + x^{12} + x^{11} + x^{11} + x^{10} + x^{9} + x^{7} + x^{6} + x^{5}$$

= $x^{13} + x^{12} + x^{10} + x^{9} + x^{7} + x^{6} + x^{5}$
= $x^{6} + x^{4} + x^{3} + x^{2} \mod x^{8} + x^{4} + x^{3} + x + 1$

Division of $x^{13} + x^{12} + x^{10} + x^9 + x^7 + x^6 + x^5$ by $x^6 + x^4 + x^3 + x^2$ is illustrated below.

	x^{13}	$+x^{12}$	$+x^{10}$	$+x^{9}$		$+x^{7}$	$+x^{6}$	$+x^{5}$			
$-x^5(x^8 + x^4 + x^3 + x^2 + 1) =$	x^{13}			$+x^{9}$	$+x^{8}$		$+x^{6}$	$+x^{5}$			
		x^{12}	$+x^{10}$		$+x^{8}$	$+x^7$					
$-x^4(x^8 + x^4 + x^3 + x^2 + 1) =$		x^{12}			$+x^{8}$	$+x^{7}$		$+x^{5}$	$+x^4$		
			x^{10}					$+x^{5}$	x^4		
$-x^2(x^8 + x^4 + x^3 + x^2 + 1) =$			x^{10}				x^6	$+x^{5}$		x^3	x^2
							x^6		$+x^{4}$	$+x^3$	$+x^{2}$

12.9.3 Hash Functions

Universal Hash Families ADD PARAGRAPH ON MOTIVATION integrate material with Chapter

Let $M = \{1, 2, ..., m\}$ and $N = \{1, 2, ..., n\}$ where $m \ge n$. A family of hash functions $H = \{h | h : M \to N\}$ is said to be 2-universal if for all x and y, $x \ne y$, and for h chosen uniformly at random from H,

$$Prob\left[h\left(x\right) = h\left(y\right)\right] \le \frac{1}{n}$$

Note that if H is the set of all possible mappings from M to N, then H is 2-universal. In fact $Prob [h (x) = h (y)] = \frac{1}{n}$. The difficulty in letting H consist of all possible functions is that a random h from H has no short representation. What we want is a small set H where each $h \in H$ has a short representation and is easy to compute.

Note that for a 2-universal H, for any two elements x and y, h(x) and h(y) behave as independent random variables. For a random f and any set X the set $\{f(x) | x \in X\}$ is a set of independent random variables.

12.9.4 Application of Mean Value Theorem

The mean value theorem states that if f(x) is continuous and differentiable on the interval [a, b], then there exists $c, a \leq c \leq b$ such that $f'(c) = \frac{f(b) - f(a)}{b-a}$. That is, at some point between a and b the derivative of f equals the slope of the line from f(a) to f(b). See Figure 12.9.4.

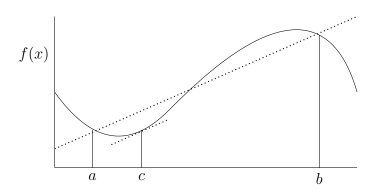


Figure 12.3: Illustration of the mean value theorem.

One application of the mean value theorem is with the Taylor expansion of a function. The Taylor expansion about the origin of f(x) is

$$f(x) = f(0) + f'(0)x + \frac{1}{2!}f''(0)x^2 + \frac{1}{3!}f'''(0)x^3 + \cdots$$
(12.3)

By the mean value theorem there exists $c, 0 \le c \le x$, such that $f'(c) = \frac{f(x) - f(0)}{x}$ or f(x) - f(0) = xf'(c). Thus

$$xf'(c) = f'(0)x + \frac{1}{2!}f''(0)x^2 + \frac{1}{3!}f'''(0)x^3 + \cdots$$

and

$$f(x) = f(0) + xf'(c).$$

One could apply the mean value theorem to f'(x) in

$$f'(x) = f'(0) + f''(0)x + \frac{1}{2!}f'''(0)x^2 + \cdots$$

Then there exists $d, 0 \le d \le x$ such that

$$xf''(d) = f''(0)x + \frac{1}{2!}f'''(0)x^2 + \cdots$$

Integrating

$$\frac{1}{2}x^{2}f''(d) = \frac{1}{2!}f''(0)x + \frac{1}{3!}f'''(0)x^{3} + \cdots$$

Substituting into Eq(12.3)

$$f(x) = f(0) + f'(0)x + \frac{1}{2}x^2f''(d).$$

12.9.5 Sperner's Lemma

Consider a triangulation of a 2-dimensional simplex. Let the vertices of the simplex be colored R, B, and G. If the vertices on each edge of the simplex are colored only with the two colors at the endpoints then the triangulation must have a triangle whose vertices are three different colors. In fact, it must have an odd number of such vertices. A generalization of the lemma to higher dimensions also holds.

Create a graph whose vertices correspond to the triangles of the triangulation plus an additional vertex corresponding to the outside region. Connect two vertices of the graph by an edge if the triangles corresponding to the two vertices share a common edge that is color R and B. The edge of the original simplex must have an odd number of such triangular edges. Thus, the outside vertex of the graph must be of odd degree. The graph must have an even number of odd degree vertices. Each odd vertex is of degree 0, 1, or 2. The vertices of odd degree, i.e. degree one, correspond to triangles which have all three colors.

12.9.6 Prüfer

Here we prove that the number of labeled trees with n vertices is n^{n-2} . By a labeled tree we mean a tree with n vertices and n distinct labels, each label assigned to one vertex.

Theorem 12.28 The number of labeled trees with n vertices is n^{n-2} .

Proof: (Prüfer sequence) There is a one-to-one correspondence between labeled trees and sequences of length n-2 of integers between 1 and n. An integer may repeat in the sequence. The number of such sequences is clearly n^{n-2} . Although each vertex of the tree has a unique integer label the corresponding sequence has repeating labels. The reason for this is that the labels in the sequence refer to interior vertices of the tree and the number of times the integer corresponding to an interior vertex occurs in the sequence is related to the degree of the vertex. Integers corresponding to leaves do not appear in the sequence.

To see the one-to-one correspondence, first convert a tree to a sequence by deleting the lowest numbered leaf. If the lowest numbered leaf is i and its parent is j, append j to the tail of the sequence. Repeating the process until only two vertices remain yields the sequence. Clearly a labeled tree gives rise to only one sequence.

It remains to show how to construct a unique tree from a sequence. The proof is by induction on n. For n = 1 or 2 the induction hypothesis is trivially true. Assume the induction hypothesis true for n - 1. Certain numbers from 1 to n do not appear in the sequence and these numbers correspond to vertices that are leaves. Let i be the lowest number not appearing in the sequence and let j be the first integer in the sequence. Then i corresponds to a leaf connected to vertex j. Delete the integer j from the sequence. By the induction hypothesis there is a unique labeled tree with integer labels $1, \ldots, i - 1, i + 1, \ldots, n$. Add the leaf i by connecting the leaf to vertex j. We need to argue that no other sequence can give rise to the same tree. Suppose some other sequence did. Then the i^{th} integer in the sequence must be j. By the induction hypothesis the sequence with j removed is unique.

Algorithm

Create leaf list - the list of labels not appearing in the Prüfer sequence. n is the length of the Prüfer list plus two. while Prüfer sequence is non empty do begin p =first integer in Prüfer sequence e =smallest label in leaf list Add edge (p, e)Delete e from leaf list Delete p from Prüfer sequence If p no longer appears in Prüfer sequence add p to leaf list end There are two vertices e and f on leaf list, add edge (e, f)

12.10 Exercises

Exercise 12.1 What is the difference between saying f(n) is $O(n^3)$ and f(n) is $o(n^3)$?

Exercise 12.2 If $f(n) \sim g(n)$ what can we say about f(n) + g(n) and f(n) - g(n)?

Exercise 12.3 What is the difference between \sim and Θ ?

Exercise 12.4 If f(n) is O(g(n)) does this imply that g(n) is $\Omega(f(n))$?

Exercise 12.5 What is $\lim_{k\to\infty} \left(\frac{k-1}{k-2}\right)^{k-2}$.

Exercise 12.6 Select a, b, and c uniformly at random from [0,1]. The probability that b < a is $\frac{1}{2}$. The probability that c < a is $\frac{1}{2}$. However, the probability that both b and c are less than a is $\frac{1}{3}$ not $\frac{1}{4}$. Why is this? Note that the six possible permutations abc, acb, bac, cab, bca, and cba, are all equally likely. Assume that a, b, and c are drawn from the interval (0,1]. Given that b < a, what is the probability that c < a?

Exercise 12.7 Let A_1, A_2, \ldots, A_n be events. Prove that $Prob(A_1 \cup A_2 \cup \cdots A_n) \leq \sum_{i=1}^n Prob(A_i)$

Exercise 12.8 Give an example of three random variables that are pairwise independent but not fully independent.

Exercise 12.9 Give examples of nonnegative valued random variables with median >> mean. Can we have median << mean?

Exercise 12.10 Consider n samples x_1, x_2, \ldots, x_n from a Gaussian distribution of mean μ and variance σ . For this distribution $m = \frac{x_1+x_2+\cdots+x_n}{n}$ is an unbiased estimator of μ . If μ is known then $\frac{1}{n} \sum_{i=1}^{n} (x_i - \mu)^2$ is an unbiased estimator of σ^2 . Prove that if we approximate μ by m, then $\frac{1}{n-1} \sum_{i=1}^{n} (x_i - m)^2$ is an unbiased estimator of σ^2 .

Exercise 12.11 Given the distribution $\frac{1}{\sqrt{2\pi^3}}e^{-\frac{1}{2}(\frac{x}{3})^2}$ what is the probability that x > 1?

Exercise 12.12 $e^{-\frac{x^2}{2}}$ has value 1 at x = 0 and drops off very fast as x increases. Suppose we wished to approximate $e^{-\frac{x^2}{2}}$ by a function f(x) where

$$f(x) = \begin{cases} 1 & |x| \le a \\ 0 & |x| > a \end{cases}$$

What value of a should we use? What is the integral of the error between f(x) and $e^{-\frac{x^2}{2}}$?

Exercise 12.13 Given two sets of red and black balls with the number of red and black balls in each set shown in the table below.

	red	black
Set 1	40	60
Set 2	50	50

Randomly draw a ball from one of the sets. Suppose that it turns out to be red. What is the probability that it was drawn from Set 1?

Exercise 12.14 Why cannot one prove an analogous type of theorem that states $p(x \le a) \le \frac{E(x)}{a}$?

Exercise 12.15 Compare the Markov and Chebyshev bounds for the following probability distributions

1. $p(x) = \begin{cases} 1 & x = 1 \\ 0 & otherwise \end{cases}$ 2. $p(x) = \begin{cases} 1/2 & 0 \le x \le 2 \\ 0 & otherwise \end{cases}$

Exercise 12.16 Let s be the sum of n independent random variables x_1, x_2, \ldots, x_n where for each i

$$x_i = \begin{cases} 0 & Prob & p \\ 1 & Prob & 1-p \end{cases}$$

- 1. How large must δ be if we wish to have $Prob(s < (1 \delta)m) < \varepsilon$?
- 2. If we wish to have $Prob(s > (1 + \delta)m) < \varepsilon$?

Exercise 12.17 What is the expected number of flips of a coin until a head is reached? Assume p is probability of a head on an individual flip. What is value if p=1/2?

Exercise 12.18 Given the joint probability

P(A,B)	A=0	A=1
B=0	1/16	1/8
B=1	1/4	9/16

- 1. What is the marginal probability of A? of B?
- 2. What is the conditional probability of B given A?

Exercise 12.19 Consider independent random variables x_1 , x_2 , and x_3 , each equal to zero with probability $\frac{1}{2}$. Let $S = x_1 + x_2 + x_3$ and let F be event that $S \in \{1, 2\}$. Conditioning on F, the variables x_1 , x_2 , and x_3 are still each zero with probability $\frac{1}{2}$. Are they still independent?

Exercise 12.20 Consider rolling two dice A and B. What is the probability that the sum S will add to nine? What is the probability that the sum will be 9 if the roll of A is 3?

Exercise 12.21 Write the generating function for the number of ways of producing chains using only pennies, nickels, and dines. In how many ways can you produce 23 cents?

Exercise 12.22 A dice has six faces, each face of the dice having one of the numbers 1 though 6. The result of a role of the dice is the integer on the top face. Consider two roles of the dice. In how many ways can an integer be the sum of two roles of the dice.

Exercise 12.23 If a(x) is the generating function for the sequence a_0, a_1, a_2, \ldots , for what sequence is a(x)(1-x) the generating function.

Exercise 12.24 How many ways can one draw n a's and b's with an even number of a's.

Exercise 12.25 Find the generating function for the recurrence $a_i = 2a_{i-1} + i$ where $a_0 = 1$.

Exercise 12.26 Find a closed form for the generating function for the infinite sequence of prefect squares 1, 4, 9, 16, 25, ...

Exercise 12.27 Given that $\frac{1}{1-x}$ is the generating function for the sequence $1, 1, \ldots$, for what sequence is $\frac{1}{1-2x}$ the generating function?

Exercise 12.28 Find a closed form for the exponential generating function for the infinite sequence of prefect squares 1, 4, 9, 16, 25, ...

Exercise 12.29 Prove that the L_2 norm of (a_1, a_2, \ldots, a_n) is less than or equal to the L_1 norm of (a_1, a_2, \ldots, a_n) .

Exercise 12.30 Prove that there exists a $y, 0 \le y \le x$, such that f(x) = f(0) + f'(y)x.

Exercise 12.31 Show that the eigenvectors of a matrix A are not a continuous function of changes to the matrix.

Exercise 12.32 What are the eigenvalues of the two graphs shown below? What does this say about using eigenvalues to determine if two graphs are isomorphic.



Exercise 12.33 Let A be the adjacency matrix of an undirected graph G. Prove that eigenvalue λ_1 of A is at least the average degree of G.

Exercise 12.34 Show that if A is a symmetric matrix and λ_1 and λ_2 are distinct eigenvalues then their corresponding eigenvectors x_1 and x_2 are orthogonal. *Hint:*

Exercise 12.35 Show that a matrix is rank k if and only if it has k nonzero eigenvalues and eigenvalue 0 of rank n-k.

Exercise 12.36 Prove that maximizing $\frac{x^TAx}{x^Tx}$ is equivalent to maximizing x^TAx subject to the condition that x be of unit length.

Exercise 12.37 Let A be a symmetric matrix with smallest eigenvalue λ_{\min} . Give a bound on the largest element of A^{-1} .

Exercise 12.38 Let A be the adjacency matrix of an n vertex clique with no self loops. Thus, each row of A is all ones except for the diagonal entry which is zero. What is the spectrum of A.

Exercise 12.39 Let A be the adjacency matrix of an undirect graph G. Prove that the eigenvalue λ_1 of A is at least the average degree of G.

Exercise 12.40 We are given the probability distribution for two random vectors x and y and we wish to stretch space to maximize the expected distance between them. Thus, we will multiply each coordinate by some quantity a_i . We restrict $\sum_{i=1}^{d} a_i^2 = d$. Thus, if we increase some coordinate by $a_i > 1$, some other coordinate must shrink. Given random vectors $x = (x_1, x_2, \ldots, x_d)$ and $y = (y_1, y_2, \ldots, y_d)$ how should we select a_i to maximize $E(|x-y|^2)$? The a_i stretch different coordinates. Assume

$$y_i = \begin{cases} 0 & \frac{1}{2} \\ 1 & \frac{1}{2} \end{cases}$$

and that x_i has some arbitrary distribution.

$$E(|x-y|^2) = E\sum_{i=1}^d \left[a_i^2 (x_i - y_i)^2\right] = \sum_{i=1}^d a_i^2 E(x_i^2 - 2x_i y_i + y_i^2)$$
$$= \sum_{i=1}^d a_i^2 E(x_i^2 - x_i + \frac{1}{2})$$

Since $E(x_i^2) = E(x_i)$ we get. Thus, weighting the coordinates has no effect assuming $\sum_{i=1}^{d} a_i^2 = 1. \quad Why \text{ is this? Since } E(y_i) = \frac{1}{2}.$ $E(|x-y|^2) \text{ is independent of the value of } x_i \text{ hence its distribution.}$ $What \text{ if } y_i = \begin{cases} 0 & \frac{3}{4} \\ 1 & \frac{1}{4} \end{cases} \text{ and } E(y_i) = \frac{1}{4}. \text{ Then}$ $E(|x-y|^2) = \sum_{i=1}^{d} a_i^2 E(x_i^2 - 2x_iy_i + y_i^2) = \sum_{i=1}^{d} a_i^2 E(x_i - \frac{1}{2}x_i + \frac{1}{4})$ $= \sum_{i=1}^{d} a_i^2 \left(\frac{1}{2}E(x_i) + \frac{1}{4}\right)$

To maximize put all weight on the coordinate of x with highest probability of one. What if we used 1-norm instead of the two norm?

$$E(|x-y|) = E\sum_{i=1}^{d} a_i |x_i - y_i| = \sum_{i=1}^{d} a_i E|x_i - y_i| = \sum_{i=1}^{d} a_i b_i$$

where $b_i = E(x_i - y_i)$. If $\sum_{i=1}^{d} a_i^2 = 1$, then to maximize let $a_i = \frac{b_i}{b}$. Taking the dot product of a and b is maximized when both are in the same direction.

Exercise 12.41 Maximize x+y subject to the constraint that $x^2 + y^2 = 1$.

Exercise 12.42 Draw a tree with 10 vertices and label each vertex with a unique integer from 1 to 10. Construct the Prfer sequence for the tree. Given the Prfer sequence recreate the tree.

Exercise 12.43 Construct the tree corresponding to the following Prfer sequences

- 1. 113663
- 2. 552833226

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