Chapter 1. The J-curve and the J distribution

“... ecologists tend to consume scientific knowledge rather than produce it.”

Whatever one might mean by the term “biodiversity,” the main thrust of all inquiries conducted under that name is the desire to say something meaningful about species of living things in their natural habitat. In particular, the population sizes of various species, whether in relative or absolute terms, become the central focus. The most comprehensive studies address whole communities, namely all the species within a particular group and living within a specific area, as might be defined by a field biologist. One might wish to know how many species of butterflies there are in a particular forest, for example, along with their relative or absolute abundances. Or one might wish to study a community of aquatic plants in a tropical embayment with the same general aim in mind. Literally thousands of such studies have been conducted for more than 100 years, in almost every clime, habitat, and living kingdom. In all cases - or nearly all cases - the investigator has had recourse to only one portal of inspection, namely samples collected from or recorded within the community in question.

The main subject of this book is the relationship of the abundances in communities (usually unknown) and the abundances that appear when samples are taken of that community. Short of a Faustian magic mirror that shows us a whole community, along with all its living components, we are stuck with samples. As indicated in the description of the frontispiece, the (so-called) J-curve is ubiquitous in samples of natural communities and there is a clear consensus to that effect, reached in the last decade. (McGill et al. 2007). The random sample of a non-natural community such as the plants in a garden or the animals in a zoo are likely to have a rather different distribution of abundances. The relationship between the two sets of abundances is described theoretically by the Pielou transform (sections 3.5 and 5.3) and is made practical in working methods for estimating abundances in a community based on one or more samples of it.

The theory proposed here to account for the ubiquitous J-curve has an associated statistical manifestation previously called the Logistic-J distribution (Dewdney 1998, 2003), but since renamed the J distribution owing to the potential for confusion with the “logistic distribution” (Balakrishnan 1991) with which it has nothing in common. Like other proposals to account for this shape, the distribution is oblivious to the actual species that have a given abundance; only treating the number of species having that abundance as relevant. The distribution is a pure hyperbola* and the corresponding “hyperbolic” theory has expanded in recent years into a mathematical tool kit for the study of populations in communities that goes beyond the estimation of abundances.

* The “hyperbolic distribution”. (Barndorff-Nielsen, 1978) is actually log-hyperbolic.

1.1. The stochastic species hypothesis and the J distribution

The theory described in this monograph begins with the stochastic species hypothesis and its logical consequent, the J distribution. In its simplest form the hypothesis is embodied in a stochastic system in which a large number of individuals are partitioned into sets called species. Individuals are chosen at random and either reproduce or die with equal probability. After each such fundamental operation the species count is either incremented or decremented, as appropriate. The hypothesis may be extended in various ways, as in Chapter 7 where it is applied to communities in the form of a binomial distribution of
population changes over extended periods of time.

As shown in Appendix A.1, a stochastic system produces a specific distribution that we will later claim, on the basis of both theoretical considerations and an extensive study of the sample literature, to manifest in the distribution of abundances both in communities and in samples of them.

The J distribution is essentially the standard hyperbola, as shown in Figure 1.1, translated in the horizontal direction by the amount $\epsilon$ (epsilon) and in the vertical direction by $\delta$ (delta). If we use the variables $x'$ and $y'$ for the primary axis system, the formula for the hyperbola is $y' = 1/x'$

Figure 1.1. The hyperbola as the basis of the J-distribution

The variables $x$ and $y$ for the displaced system obey the following equalities,

$$x = x' - \epsilon$$
$$y = y' - \delta,$$

and the formula for the probability density function (pdf) becomes

$$f(x) = c(1/(x+\epsilon) - \delta); \quad 0 \leq x \leq \Delta - \epsilon,$$

$$= 0; \quad x \geq \Delta - \epsilon$$

where $\Delta = 1/\delta$ and $c$ is a normalizing constant (a function of $\epsilon$ and $\delta$) that gives the pdf an area of unity, as required by all probability density functions. This “density” represents the probability of finding a species with an abundance represented by an infinitesimal interval on the abundance axis. The constant $c$ has the following formula:
\[ c = (\ln((\Delta/\varepsilon) - 1 + \varepsilon/\Delta)^{-1} \]

In most cases, one may neglect the term \( \varepsilon/\Delta \), which tends to be rather tiny.

It is interesting that the density function just described is symmetrical under inversion. If one omits the constant \( c \) for a moment and writes,

\[ y = 1/(x+\varepsilon) - \delta, \]

one may express \( y \) as a function of \( x \) by inverting the function, i.e.,

\[ x = 1/(y+\delta) - \varepsilon. \]

The algebraic symmetry merely reflects the geometric symmetry of the hyperbolic function about the diagonal \( y' = x' \).

In any case the J distribution is a two-parameter distribution. The inverses of the constants \( \varepsilon \) and \( \delta \), namely \( \varepsilon' \) and \( \delta' \), represent the species peak and the logistic limit, respectively. The first parameter \( \varepsilon' \) (when multiplied by \( Rc \), the number of species and the constant \( c \)) bounds the number of species in the lowest abundance category, as well as all others. The second parameter \( \delta' \) bounds the abundance of the largest population. Both “bounds” operate statistically, being simply averages of their respective data. In Appendix A.1 the following elements of the J distribution are placed on a sound mathematical footing as implications of the stochastic species hypothesis:

1. The hyperbolic shape of the distribution is derived in Theorem A.3, Section A.1.5
2. The parameter \( \delta \) emerges in theorem A.4, Section A.1.5
3. The parameter \( \varepsilon \) emerges in Section A.1.6

1.2. Appearance of the J distribution

The J distribution first appeared with certainty from a computer simulation written in the mid-1990s to simulate a mutually predatory community of protists. As a longtime sampler of local waters (Dewdney 1996, 2010), I had plenty of microbial data on hand. As a longtime teacher of stochastic simulation and statistics, I had the tools and background necessary to the project.

When it first occurred to me to write such a program, I had already been observing the J-curve in plots of my abundance data, but I had expected something more like a normal curve, with a concentration of species about some mean abundance. Although I was well aware that no microbial community could possibly consist of mutually predatory species of, say, ciliates, it seemed at the time that if any such “community” could produce a unimodal, normal distribution of abundances, it would be this mutually predatory one.

A simulation clock governed the flow of events in the system; at each tick of the clock two individual “organisms” would be selected from the total community population. Without having to represent the
organisms at all, the simulation would simply ensure that the species of the first individual had its population incremented by 1, the species-token moving, in consequence, one unit to the right along the abundance axis. Meanwhile the species of the second individual would have its population decremented, with its species shifting one position to the left. In such a system, the total number of individuals would be preserved by the “predation” operation.

The original purpose of the simulation was to demonstrate that such a community would achieve a balance of populations, with all species fluctuating about the same mean population size. This did not happen. Indeed, a J-shaped curve stubbornly emerged every time the simulation was run! Each run would begin with a single spike, with all populations occupying the same abundance category. Over time, the spike would flatten into a skewed, bell-shaped curve, where I had expected the process essentially to stabilize. Instead the process continued as the curve continued to flatten out, even as some species piled up at the low end of the abundance scale while others fled randomly to the high end. In most of these mini-experiments, I kept the average abundance low in order to encourage a more definite shape. Accordingly, I kept the extirpation switch in the “off” position in order to allow the shape to build. (Descriptions of this and other systems used in this research will be found in Appendix A.3.)

During the equilibrium phase of the simulation, histograms with a low mean abundance bore a close similarity to the field samples that I had already begun to collect from biosurveys of other kinds of communities. Indeed, they were (visually at least) indistinguishable from them. Yet only a small fraction of the samples I had been examining involved predators of any kind. It took a surprising amount of time (in retrospect) to realize that the simulation was not about predation, with one organism dying while another reproduced in consequence. It was about the birth/death processes itself, in particular the equality of probabilities of births versus deaths. After all, there was no actual predation in the dynamical system. Whenever one species moved to the right, some other species would move to the left. It was a clear case of a specific model turning out to be rather general, after all.

An initial result that appeared to confirm the presence of a hyperbolic function relied on the analysis of a slightly more general system in which a random individual was selected from the set of all individuals and either duplicated or deleted, both events taking place with equal probability. (See Appendix A.1 for the Equilibrium Theorem)

After a year of studying the more general system, I obtained a more exact result: the equilibrium solution of the system was indeed a hyperbola, but translated downward by a small amount that I decided to call “delta.” The formula was

\[ k(1/x - \delta), \]

where \( k \) is a constant (Dewdney 1998a). (See Appendix A.1 for the Logistic Theorem.) New simulation programs were written to explore the idea of equiprobable birth/death processes under varying conditions on the balance of probabilities governing them. The probability of birth, for example, could exceed that of death for a time, but it would eventually move in the other direction -- all at random. These more general versions of the dynamical system produced the same J-shaped curves as the earlier system. Indeed, they were indistinguishable from the histograms of field surveys that I had already been to collecting. One could hardly be blamed for suspecting that the J-curve was a universal phenomenon.
In the new simulations, each individual “organism” had an approximately equal probability of dying or reproducing at each moment. Populations fluctuated randomly but ultimately a few became larger, while many became smaller. The result, as in the case of the predation system, was distinctly counterintuituitive and illustrates the dangers of armchair theorizing. At equilibrium, which never failed to develop, the histogram would be frozen and compared with a hyperbola (visually). The fit was often rather good, taking into account normal statistical fluctuations. Since the net effect of the births and deaths in this program was to preserve the total “biomass” of the system, the logistic limit could be applied and the same formula derived from the predation system applied here, as well. At this point a general mechanism suggested itself. Called the stochastic species hypothesis, it postulated equal probabilities of birth and death over unit time periods, with corresponding changes in population thus expected.

Following an intensive but disappointing literature review of other species-abundance models, a massive study was launched. With the aid of graduate students in our Department of Biology, I began a random collection of biosurvey papers, ensuring that all the major classes of biota were covered as they went. Each of the resulting histograms was compared with a version of the J distribution that shared the same mean and height of initial peak; given values for these quantities, the values of the parameters $\varepsilon$ and $\Delta$ are readily determined. (See Transfer Equations in Appendix A.1.) The resulting chi square scores were normalized to 10 degrees of freedom to make cross-comparisons possible. (See the scores histogram in Figure 1.2) The average score that emerged from the study was 10.43, very close to the optimum score of 10.0 and well separated from the average score that emerged from parallel tests of the distribution that most closely resembled the hyperbolic curve. The log-series distribution mentioned in the previous section scored over 13. It was unnecessary to test other distributions, since they resembled the J distribution even less. Only something with a very similar shape could hope to score as well (Dewdney 2003). In short, there is no “room” for another distribution in view of the optimality of the scores. The results of this metastudy amounted to what can only be called “strong support” of the stochastic species hypothesis and therefore of the J distribution that emerges from it as a universal natural phenomenon.

1.3 The community and the sample

Figure 1.1 illustrates the relationship between a natural community and a sample of it. Each little square represents a species, its position on the horizontal axis representing its abundance. It will be a general feature of the histograms shown in this book that the numerical label associated with each abundance appears at the right hand side of the corresponding interval on the abundance axis. This is a technical device that harmonizes discrete and continuous versions of the hyperbolic distribution. Some theoretical abundances, as well as some empirical ones, are fractional, with values such as 3.2, 3.7, and 3.8, for example, all gathered into the category 4.0. In general, the kth category would embrace all abundances lying in the half-open interval, $(k-1, k]$.

It will be a common experience for readers of this book to encounter irregular histograms like the ones in Figure 1.1 where, due to horizontal space limitations, not all species are shown. In cases where it matters, the missing abundances are always given. In cases where it doesn’t, the reader may imagine them as being present, nevertheless. Indeed, the J distribution indicates just how far out on the abundance axis such species might be expected to occur.
Figure 1.2  The relationship between abundances in a community and those of a sample

The histograms shown in Figure 1.2 are imaginary, but reflect what one might call realistic variability. Those with experience of actual sample histograms may well accept the sample data as typical, but who has seen the histogram of an entire community? The community histogram is simply an indication of what a community might look like, given the nature of the sampling process. The horrifying sight presented in the sample histogram of so many species seemingly about to be extirpated vanishes at the sight of the community histogram with such a peak being absent or weakly developed. In any case, from the point of view of the present theory, the initial peaks are far from “anomalous” (Coddington et al., 2009). As this monograph shows, they are entirely natural. It must be remembered that, in general, the abundance of a species in a sample is usually much lower than its abundance in the community from which the sample was taken. For example, a species of abundance 1 in the sample histogram might well have abundance 27 in the source community. The sampling process has “mapped” the higher community abundance into one of relatively low abundance.

Although the focus of this work is ultimately the community under investigation, we have only samples to work with. As a result, most of the histograms presented in this book belong to either hypothetical or actual samples. As illustrated in the frontispiece, they are generally characterized by a high initial peak that is followed by a rapid descent in bar height, whether smooth or ragged. The descent becomes more gradual with ever higher abundances, there being ever fewer species at these abundances. The theoretical curves that represent these histograms have the same general shape, although they have a smoother appearance.

The following sections in this introductory chapter include a brief review of other proposals for abundance distributions, along with an explanation of goodness of fit tests and their use in both negative and positive mode. An introduction to the J-distribution begins with the probability density function (pdf), an explanation of the two associated parameters and a formula for the mean $\mu$. The chapter ends with a brief history of the author’s early work in this area, providing another doorway to understanding.
1.4 The need for an appropriate theory

It is an unusual development in any science that a plethora of formulas should result from the attempt to capture a natural phenomenon in mathematical terms. In the normal course of research any formula found wanting would be discarded and a new formula developed. Since the early 1940s about a dozen formulas for species abundances (in samples) have been proposed and none of them discarded. Here are the principal ones discussed by Magurran (1988). She also mentions another three proposals, for a total of seven.

1. The geometric series (May 1975)
2. The log-series distribution (Fisher, Corbett & Williams 1943)
3. The lognormal distribution (Preston 1948), (See Sections 3.6 and 3.6.1)
4. The broken stick model (MacArthur 1957)

Among the other proposals discussed by Magurran, special mention should be made of the dynamical model of Hughes (1986), the only proposal prior to the 1990s to be tested against an appropriately large number of samples from the field. Unfortunately, Hughes was never able to find a closed formula for the J-shaped curves that his data produced.

It is difficult to characterize succinctly, without committing the sin of oversimplification, how ecologists interested in theory have worked in the past. However, it is fair to say that the various proposals just indicated have been compared with various sets of field data over the years with the aim of discovering which proposals fit the field data best. This seems like a perfectly logical procedure, but it contains a very dangerous pitfall in the form of an unstated (and apparently unrealized) assumption. From the point of view of statistical theory, the outcome of this method is (when one thinks about it) predictable, with each distribution fitting best with certain data sets and not fitting others as well. For example, four of the proposals listed above are described by Magurran, in her research summary, as follows. Based on the rank-abundance representation of field data (Section 4.2 of the present work), the geometric series best fitted (apparently by visual inspection) plant data from a sub-alpine forest, the log-series and the lognormal best fitted a histogram of plant data from a deciduous forest, and the broken stick model best fit a survey of nesting birds in a deciduous forest.

The methodology of comparing a handful of field histograms with various theoretical distributions has persisted into the new millennium. For example, Hubbell (2001) continues this methodology and, in a sense, brings it to a climax, drawing support for various manifestations of his zero-sum multinomial distribution from seven studies of various biotic groups, from trees to birds and, not surprisingly, finds a good (visual) fit within a milieu that contains previously proposed distributions implicitly. Connolly et al (2005) compared four different distributions with survey data involving corals and fish in Pacific Ocean reefs, finding that some distributions fit the data from some reefs better than others, where other distributions did better.

As more and more data sets are compared with the limited number of distribution proposals under active consideration, it is not surprising that each camp builds up followers, so to speak. A summary of the situation provided by Diserud and Engen (2000) points out that “A large number of data sets from ecological communities correspond well to the model in which the abundances are lognormally distributed . . . ”, later pointing out in the same breath that “lognormal models may often provide a rather
bad fit to observed data.”

As if it were a tacit admission that something is wrong, there has been a long tradition in theoretical ecology of attempting to unify somehow the various distribution proposals into one. It is not difficult to find mathematical generalizations of the proposals mentioned above. But was it the unsatisfactory results of such unification projects that has spawned a cottage industry of attempts? Beginning with Sugihara (1980) and culminating with (Deserud and Engen 2000), then (Hubbell 2001), the unification process has served to keep alive the idea that each proposal has its niche, so to speak. More recently, a group of 18 authors (McGill, Etienne, et al. 2007) have proposed “moving beyond single prediction theories to integration within an ecological frame-work.” The article demonstrates clearly a continuing confusion about the roles of species abundance distributions in an ecological context. It makes little sense to “move beyond” the present situation until it is corrected.

The underlying methodological weakness has resulted in no little soul-searching, as when a year later the 14-author paper by Doak, Estes et al. (2008) declares that, in view of science being full of surprises, “... it is not so surprising, so to speak, that we frequently face outcomes of experiments and observations that leave us scratching our heads, wondering how we could have been so wrong in our expectations.” In the context of the methodological problems just described, it isn’t surprising at all.

Two main assumptions appear to support the traditional approach to the assessment of proposals. It isn’t clear from my reading of the literature that either assumption has ever been explicitly stated in some journal article, but most would agree that they would be necessary assumptions for the method just described to work:

a) the shape of the sample histogram, however its abundances are plotted, reflects in a specific way the shape of the community.

b) the shape of a community histogram is stable in the sense that two samples that are well separated in time or space will reflect that community.

The first assumption is an article of faith with field biologists. If the samples they so painstakingly gathered in the wild turned out to have no such relation with the community, there could only be despair. In fact, the distribution underlying the sample “reflects” the distribution that underlies the community in a statistical manner. The underlying distributions are the same, on average, but for a change in parameter values. (Dewdney 2002) Thus, whatever one means by the (deliberately vague) word “reflect”, the first assumption is completely solid.

A reasonable abstract description of any “community” can be framed in the context of the upper histogram in Figure 1.2. As I have pointed out, the histogram of a community would have a rather strung-out appearance, in comparison with the histogram of a sample of it. Over time, the various populations that make up the community will all fluctuate, some increasing, some decreasing, with occasional changes of direction, seemingly at random, among all of the populations. In terms of the histogram itself, one may visualize these fluctuations as a sort of vibration when the film is speeded up, so to speak. Species at the high abundance end vibrate at a higher frequently because their populations are much larger, with births and deaths occurring at a higher rate over time. At the low abundance end, in corresponding fashion, the
species move in more sedate fashion, slowing rather quickly as they approach extirpation.

The reasons for the apparent randomness in the motion of these species are typically myriad. Suffice it to say that the overall shape of the histogram, however one described it, would also change accordingly. All proposals for the underlying theoretical distribution share what might be called the J-attribute (albeit in a mild form): an initial peak is followed by a smooth decline. All samples from the field also share the J-attribute, albeit more strongly peaked and with a steeper initial decline. Within such a descriptive framework one may find room for any of the extant theoretical proposals. Samples of the putative community might well, over time, exhibit the full range of these shapes. Would they change enough to defy a description as “geometric”, “log-series”, “log-normal”, or “broken stick”? The answer is simple: It would change its overall shape enough, over time, to defy any proposal, including the centerpiece of this book, the J distribution.

I cannot account for the reasoning behind Assumption b), but it may have something to do with the notion that large sample size guarantees statistical significance. In other words, the large size of the field samples that have been used to promote one distribution over another were themselves thought to guarantee that the overall results were somehow determinative.

According to the first assumption, as a community distribution changes its shape over time, so would the samples taken of it. Studies that would attach significance to such changes (Magurran 2007) fall into essentially the same trap. Changes in the shape of a community distribution will happen willy-nilly and the new variations of the J-curve, as revealed (or not) by the sample distribution have little actual significance. Hyperbolic theory tells us that living communities in general have no particular shape beyond displaying the J-attribute. The relationship between communities and samples of them is explored fully in Chapter 3.

The sub-alpine community of forest plants that supported the geometric distribution today, may well betray it utterly 50 years later, favoring the log-series distribution instead, perhaps. With this view of natural communities in mind, none of the conclusions about proposed distributions reached by this method can be accepted as anything but coincidences. Made 30 years earlier on corresponding data, the order of the four distributions might well be scrambled. In view of this unfortunate methodology, one could substitute any four distributions one liked -- made up for the occasion -- then rewind the historical tape and watch very much the same papers appear, favoring one distribution or another or perhaps attempting to unify them! To put the same point more precisely, it is certainly possible to submit all 125 datasets in the metastudy reported in Chapter 8 to a simple test. It would be a bit labor-intensive, but one could compare all 125 datasets with each of the theoretical distributions that have appeared in the literature to date. It is an elementary observation that every proposal will have some datasets fit that distribution better than any other. Some will have more than others, of course, and I think it likely that the J-distribution would have the most.

In the remainder of this chapter and in the chapters to come, I will argue that there can be only one universal statistical descriptor of abundances in natural communities.

As for other problems with the methodology just described, one might also point out that goodness-of-fit tests that are standard in other fields were rarely employed in the assessment of these shapes, with only two having shown up in articles. Tests such as the Chi square or Kolmogorov-Smirnov take two histograms and
compare them, producing a numerical measure of the degree of the similarity. In a field that yearns for quantitative precision, why would anyone neglect a perfectly good numerical measure of similarity between theoretical and empirical histograms? The fact that such measures of similarity have not been used may explain why there is so little awareness among theorists of the enormous variation to be expected from a single underlying distribution. The chi square curve (See Figure 1.2) has a long tail to tell, so to speak, of the many field histograms that will fit rather poorly without failing to arise from the distribution under test.

The effort to fit individual field histograms to specific theoretical distributions reminds one of the story of the child who was found one day with three buckets into which he was busily sorting pennies. The buckets were labeled “unlucky” “normal” and “lucky”. He would take a penny from a pile beside him and flip it 10 times, counting the number of heads, then placing the penny in the “lucky” bucket if it came up heads more than seven times. If it came up heads fewer than four times, he would deposit it in the “unlucky” bucket. The remainder ended up in the bucket labeled “normal”. The specific biotic source of a sample no more inheres in a single field histogram than the element of luck inheres in a penny. In this little tale the role of the overall average (p = 0.5) is played by the J distribution.

Given the difficulties just described, it may be suspected that the field of theoretical ecology is in the grip of an unacknowledged crisis. The failure to appreciate the statistical behavior of abundances in communities has become a nursery of untested proposals. The crisis is only made worse by the adoption by some ecologists of a social constructivist ethic, as enunciated by Hilborn and Mangel in *The Ecological Detective* (1997). In this milieu, “. . . there is no correct model.” Ironically, the statement was largely true up to the time of that book’s publication. The notion that all “models” are more or less acceptable is nevertheless unhelpful. At this point, the quote from Sir Francis Bacon that begins this chapter comes into play. If, in a given field of inquiry, it is not possible to be in error, then it is not possible for that field to be a science. After all, that’s exactly what the quote means.

1.3. A positive test using multi-sample data

As normally applied, goodness-of-fit tests are used to reject hypotheses. As such, the rejection, when supported by the test, applies to each sample tested. Non-rejection of a particular fit does not amount to “acceptance”, as such, since the tests are not symmetrical in this respect. Non-rejection, when based on a single sample or just a few, may be taken as evidence in favour of the null hypothesis, but as “evidence” it is weak and cannot be used, by itself, to establish anything. Since such tests are used here in affirmative rather than rejective mode, a great many samples, rather than just a few, are needed to establish the presence of an underlying distribution.

This last point is important enough to expand upon so that the reasons for the multi-test requirement are made plain. As normally used in curve-fitting applications, a test such as the chi square compiles the chi square statistic or “score” as follows.

$$\chi^2 \text{ score} = \sum \frac{(t_i - e_i)^2}{t_i}$$

The index i ranges through abundance classes 1, 2, 3, etc., and the variables t_i and e_i represent the number of species having abundance i in the theoretical and empirical distributions, respectively. The greater the
difference between the theoretical prediction ($t_i$) and the empirical datum ($e_i$), the greater is the contribution to the statistic above. Thus higher chi square scores reflect a rather poor fit to the data, while lower scores represents better fits. The same thing is true of the Kolmogoroff-Smirnov test.

Used in the normal way (Hays & Winkler 1971) a goodness-of-fit test will determine the likelihood that a given set of empirical data does not have a particular theoretical distribution. In such a case a chi square score, for example, can be used to reject the distribution if it exceeds a threshold or critical value, as given in a standard chi square table. (See Chapter 4.) In such a case, the null hypothesis (that the empirical data follow a particular theoretical distribution) is rejected. In normal parlance the hypothesis is said to be “accepted” if the score is at or below the appropriate critical value. As already pointed out, such terminology does not actually mean that the data in question have the distribution under test. After all, there are infinitely many curves, each with a different mathematical formula, that would fit the data as well or better than the distribution under test. How could a particular form be confirmed as the underlying distribution?

![Figure 1.2. Distribution of chi square scores: bar height = number of scores](image)

To use a goodness of fit test in confirmatory mode, one must have a great many datasets (say 100) and one must perform a goodness-of-fit test on all of them, compiling the scores themselves into a histogram that may be compared directly with the chi square distribution itself, as in Figure 1.2. The figure shows the chi square distribution (smooth curve) with the histogram of 125 test scores superimposed upon it. The scores are the result of the metastudy reported in Chapter 7 of this monograph. The height of bar above a given numerical category represents the number of chi square scores that fell within that category. Thus four scores with values in the range (4.00, 5.00] happened to fall in category 5.

The theoretical chi square curve clearly reflects the expectation that some samples will have very poor (i.e., large) scores, while others have very good ones. Indeed the actual scores thus achieved largely fulfill this prediction. It is a key feature of chi square theory that if a great many such scores emerge from a single underlying distribution, their average value will equal the number of degrees of freedom of the tests.
themselves. If all the tests were conducted at 10 degrees of freedom, for example, the average expected score would be 10.0 or very close to it. The chi square envelope in the figure is based on 10 degrees of freedom, while scores emerging from the metastudy were normalized to 10 degrees of freedom. The resulting average score of 10.43 is slightly greater than 10, the score bars being collectively shifted slightly to the right. In any event, it is not possible for results to be shifted to the left by any significant amount, since this would violate Pearson’s theorem. The overall fits were optimal in this sense. If other proposed distributions have test scores that are far enough away from the optimal mean to be statistically separable from it, the J distribution is clearly preferable (Dewdney 2000).

Among the other proposals, the distribution closest to the J distribution in overall shape is the log-series distribution. When put through the the same series of tests against empirical data, the log-series had a significantly higher score of 13.56, overall. It’s chi square “envelope” would overlap the one shown in Figure 1.2, but it would be displaced a few categories to the right. Other distributions proposed for the role of species/abundance descriptors, particularly the lognormal distribution, resemble the Hyperbolic far less and one does not need to subject them to the same test since their scores would be far too high to be in the running. To do as well as the J distribution, a proposed alternative must resemble it very closely. I will return to the metastudy in Chapter 8, providing all the detail necessary for its evaluation as a research tool.

It is an interesting historical fact that the British biologist C. B. Williams thought that the “hollow curves” he was seeing in lepidopteran light-trap data collectively resembled hyperbolas. (Williams 1964). He appears to have been right. His statistical colleague, R. A. Fisher, talked Williams out of the hyperbola, declaring that it was unsuitable for use as a statistical distribution, since it had a nonfinite area under it. Fisher was also right, but the idea of a translated hyperbola truncated by its axes obviously never occurred to him. Given that the subtleties of randomly fluctuating populations would not become apparent until computers were widely available, Fisher can hardly be blamed for this. (See next section.) In any event, it was not Fisher who developed goodness-of-fit tests, but his arch-rival (statistically speaking), Carl Pearson. Fisher undoubtedly knew about goodness-of-fit tests, but seems to have avoided them for some reason. Ironically, in spite of Williams’ insight, it may have been here that the pattern of inappropriate testing in theoretical ecology first developed.

The foregoing provides some background to the development of the J distribution that may be useful in clarifying its role in species-abundance studies. In the next chapter I will provide more of the mathematical details connected with the J distribution and in subsequent chapters I will show how the distribution may be used in the field to make reliable statistical inferences about communities from their samples.

1.6. Fluctuating populations

The stochastic species hypothesis (see section 7.1.1 for a full description) might seem, at first sight, to justify a great throwing up of the hands: if it’s all random, what’s the point of trying to tease out ecological mechanisms to account for population changes? The answer is simply that the stochastic species hypothesis provides a framework of understanding within which such studies may be pursued. Populations fluctuate for a huge variety of reasons, each with its own mechanism. The net effect of these mechanisms is effectively random, even though one mechanism may dominate during one period of time, another later. Weather is the ultimate determinant of many of these mechanisms and the weather itself is known to be subject to the grand mechanism of chaos (Gleick 1985) which itself produces effectively random events.
Randomness in natural populations has long been suspected, but has faced an uphill struggle in the academic forum against deterministic ideas which can be classified into a continuum. The first view is that all species maintain more or less fixed populations (Marsh 1865), a common 19th century understanding of nature. Amazingly, this view persisted to some degree throughout most of the 20th Century. For example, in a book on population genetics Wallace (1981) declares more than 100 years later, “The second thing we can say about populations is that, despite temporal fluctuations, in the long run they tend to remain constant in size.”

This “common understanding” was replaced by the notion of predator-prey cycles, as predicted by the Lotke-Volterra equations (Leigh 1968). Backed by famous datasets such as the Hudson’s Bay trapping data (Odum, 1953), the theory gained wide acceptance and is still believed by many ecologists today, in spite of experimental evidence that such cycles don’t always develop, e.g. the Gause experiments (Gause 1934), as described in (Botkin 1990). However, a more modern view, that of “density regulated” populations, held that populations did, indeed, fluctuate randomly, but only within limits imposed by the density (relative abundance) of a species (Smith 1935). Despite the extensive literature on density-regulation that has developed since Smith’s time, this concept of population behavior is not universally accepted. Thanks in part to the popularity of chaotic population dynamics (May 1974) and in part to the failure of the population-regulation school to come to firm conclusions (Cappuccino and Price 1995), there would appear to be a growing suspicion that fluctuations in natural populations are indeed random, the same view pursued by Hubbell (2001). An interesting example of accommodation between the two views addresses the concept of “density-vague” behavior in which populations are bounded away only from extreme density and extirpation (Strong, 1986). Not surprisingly, the literature is sprinkled with negative results, as in the comparison of a great many population datasets with random walks that yielded very little difference. (den Boer, 1991).

About the problems of coming to grips with population fluctuations, Botkin (1990) writes:

At the heart of the issues are ideas of stability, constancy, and balance, ideas intimately entwined in theories about nature. Perhaps one reason that the deficiencies of the theories were not examined or tested adequately by observation in the field -- out in nature -- was that ecologists were typically uncomfortable with theory and theoreticians. Doing science and creating theory were commonly distinguished as separate activities. Although theory was typically considered not to be necessary or important to the practicing ecologist, . . . theory played a dominant role in shaping the very character of inquiry and conclusions about populations and ecosystems (i.e., about nature). As Kenneth Watt wrote in 1962, ecologists had tended to believe that their science had lacked theory, while in fact it had ‘too much’ theory -- in the sense that the theory had been utilized and was influential even though it was not carefully connected to observations. ‘Field ecologists,’ those making measurements and observations in the forest and field, generally did not understand the mathematics of the logistic and of the Lotka-Volterra equations. But since physicists and mathematicians had the highest status among scientists, and since what physicists and mathematicians generally said was generally right, field ecologists tended to regard the logistic and the Lotka-Volterra equations as true. Lacking the understanding to analyse and thereby to criticize these equations, they accepted them on the basis of
authority.

The J distribution breaks this deadlock by implying that notions of stability are best applied at the community level and not at the level of populations. Species are constrained only at the high abundance end by logistic influences embedded in the myriad interactions among species within a community, between communities, and between species and the physical environment.