Indices of diversity and evenness*

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Abstract
An overview is given of the different indices used, since their introduction in the 60's, for the determination of diversity in biological samples and communities. The most commonly used indices are based on the estimation of relative abundance of species in samples. Relative abundance can also be used for either a graphical or a mathematical representation of species-abundance relationships, from which diversity indices can be deduced as well. Most common in the literature are indices either describing the richness or species number and the evenness or partitioning of individuals over species or a combination of both. The most commonly used diversity indices can be grouped in a coherent system of diversity numbers developed by Hill (1973) that includes species richness, the Simpson index and a derivation of the Shannon-Wiener index as special cases. In this system species are different only when their abundance is different. Therefore, during the last decade a number of indices have been developed that take into account the taxonomic position, trophic status or body size of the species. There is as yet no consensus as to the use of evenness indices. We apply the condition that evenness should be independent of species richness (Heip, 1974). The number of potential evenness indices is then strongly reduced. It is


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argued that the calculation of diversity or evenness indices should simply serve as descriptors of community structure and be complemented with information on ecological functioning.

Indices de diversité et régularité

Mots clés : diversité, richesse spécifique, régularité, relations espèce-abondance

Résumé
Les différents indices servant à déterminer la diversité dans les échantillons biologiques et les communautés sont passés en revue, depuis leur introduction dans les années 1960. Les indices les plus souvent employés se basent sur l’estimation de l’abondance relative des espèces dans les échantillons. L’abondance relative peut également être utilisée pour la représentation graphique ou mathématique des relations espèce-abondance, desquelles peuvent être déduits les indices de diversité. Les indices les plus souvent rencontrés dans la littérature sont ceux qui soit décrivent la richesse ou le nombre d’espèces, soit la régularité ou le regroupement des individus dans les espèces, ou bien une combinaison des deux. Les indices les plus utilisés peuvent être regroupés dans un système cohérent développé par Hill (1973), qui inclut la richesse spécifique, l’indice de Simpson et un dérivé de l’indice de Shannon-Wiener pour des cas particuliers. Selon ce système, les espèces sont différentes seulement quand leur abondance est différente. Cependant, au cours des dix dernières années, ont été développées un certain nombre d’indices qui prennent en considération la position taxonomique, le statut trophique ou bien la taille corporelle des espèces. Il n’y a pas encore de consensus sur l’utilisation des indices de régularité. Nous appliquons les conditions où la régularité devrait être indépendante de la richesse spécifique (Heip, 1974). Le nombre d’indices de régularité potentiel est alors fortement réduit. Il est démontré que le calcul de la diversité ou des indices de régularité devrait seulement servir en tant que descripteurs de la structure d’une communauté et être complété par des informations sur le fonctionnement écologique.

1. Introduction

It is common practice among ecologists to complete the description of a community by one or two numbers expressing the “diversity” or the “evenness” of the community. For this purpose a bewildering diversity of indices have been proposed and a small subset of those have become popular and are now widely used, often without much statistical consideration or theoretical justification. The theoretical developments on the use of diversity indices have been mostly discussed in the 60’s and 70’s. Although the subject continues to be debated
to this day, by the 90's their popularity in theoretical ecological work had declined. In contrast to this loss of interest from theoretical ecologists, diversity indices have become part of the standard methodology in many applied fields of ecology, such as pollution and other impact studies. They have entered environmental legislation and are again attracting attention at the turn of the century because of the surge of interest in biodiversity and the never ending quest for indicators of the status of the environment.

The basic idea of a diversity index is to obtain a quantitative estimate of biological variability that can be used to compare biological entities, composed of discrete components, in space or in time. In conformity with the "political" definition of biodiversity, these entities may be gene pools, species communities or landscapes, composed of genes, species and habitats respectively. In practice, however, diversity indices have been applied mostly to collections or communities of species or other taxonomic units. When this is the case, two different aspects are generally accepted to contribute to the intuitive concept of diversity of a community: species richness and evenness (following the terminology of Peet, 1974). Species richness is a measure of the total number of species in the community (but note already that the actual number of species in the community is usually unmeasurable). Evenness expresses how evenly the individuals in the community are distributed over the different species. Some indices, called heterogeneity indices by Peet (1974), incorporate both aspects, but HELP (1974) made the point that in order to be useful an evenness index should be independent of a measure of species richness.

Because comparison is often an essential goal, a diversity index should in principle fulfil the conditions that allow for a valid statistical treatment of the data, using methods such as ANOVA. This also requires that estimates from samples can be extrapolated to values for the statistical population. The statistical behaviour of the various indices that have been proposed is therefore a point of great importance. In practice a number of problems arise such as the definition of the entities of variability (which species, life-stages or size classes within species, functional groups, etc.?), problems of delimitation of communities and habitats, sample size, etc. that, when not accounted for, prevent the correct application of univariate statistics.

The major starting point for nearly all computations is a matrix containing stations as columns and species as rows and of which the entries are mostly abundance or biomass data. Diversity indices are univariate (and therefore do not contain all the information present in the species x stations matrix), but the same matrix can be used as the starting point for either a univariate (using other summary statistics) or a multivariate analysis. In modern ecological

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practice diversity indices are therefore nearly always used in conjunction with multivariate analyses.

The main problem in obtaining estimates of diversity is the basic but often painstaking effort necessary to collect the samples in the field and to sort, weigh and determine the organisms present in the sample. The cost and effort of the calculations are now minor in comparison. The general availability of large computing power and the wide array of easily available software has made the computations that were tedious only twenty years ago extremely easy now. The danger is that often the conditions for application of the software, e.g. checking the assumptions and conditions required for using specific statistical tests against the characteristics of the available data, are not considered.

2. Species-abundance distributions

2.1. Introduction

Nearly all diversity and evenness indices are based on the relative abundance of species, i.e. on estimates of \( p_i \) in which

\[
p_i = \frac{N_i}{N}
\]

with \( N_i \) the abundance of the \( i \)-th species in the sample, and

\[
N = \sum_{i=1}^{S} N_i
\]

with \( S \) the total number of species in the sample.

If one records the abundances of different species in a sample (and estimates them in a community), it is invariably found that some species are rare, whereas others are more abundant. This feature of ecological communities is found independent of the taxonomic group or the area investigated. An important goal of ecology is to describe these consistent patterns in different communities, and explain them in terms of interactions with the biotic and abiotic environment.

One can define a "community" as the total set of organisms in an ecological unit (biotope), but the definition used must always be specified as to the actual situation that is investigated. There exist no entities in the biosphere with absolutely closed boundaries, i.e. without interactions with other parts. Therefore, some kind of arbitrary boundaries must always be drawn. PIELOU (1975) recommends that the following features should be specified explicitly:

1. the spatial boundaries of the area or volume containing the community and the sampling methods;
2. the time limits between which observations were made;
3. the set of species or the taxocene (i.e. the set of species belonging to the same taxon) treated as constituting (or representing) the community.

The results of a sampling program of the community come as species lists, indicating for each species a measure of its abundance (usually number of individuals per unit surface, volume or catch effort, although other measures, such as biomass, are possible). Many methods are used to plot such data. The method chosen often depends on the kind of model one wishes to fit to the data. Different plots of the same (hypothetical) data set are shown in figure 1.

It can readily be seen that a bewildering variety of plots is used. They yield quite different visual pictures, although they all represent the same data set. Figures 1A–D are variants of the Ranked Species Abundance (RSA) curves. The S species are ranked from 1 (most abundant) to S (least abundant). Density (often transformed to percentage of the total number of individuals N) is plotted against species rank. Both axes may be on logarithmic scales. It is especially interesting to use a log-scale for the Y-axis, since then the same units on the Y-axis may be used to plot percentages and absolute numbers (there is only a vertical translation of the plot).

In so-called “k-dominance” curves (LAMBSHEAD et al., 1983) (figure 2A–D), the cumulative percentage (i.e. the percentage of total abundance made up by the k-th most dominant plus all more dominant species) is plotted against species rank k or log rank k. To facilitate comparison between communities with different numbers of species S, a “Lorenzen curve” may be plotted. Here the species rank k is transformed to \((kjS) \times 100\). Thus the X-axis always ranges between 0 and 100 (figure 2C).

The “collector’s curve” (figure 2D) addresses a different problem. When one increases the sampling effort, and thus the number of animals N caught, new species will appear in the collection. A collector’s curve expresses the number of species as a function of the number of specimens caught. Collector’s curves tend to flatten out as more specimens are caught. However, due to the vague boundaries of ecological communities they often do not reach an asymptotic value: as sampling effort (and area) is increased, so is the number of slightly differing patches.

The plots in figure 3A–D are species-abundance distributions. They can only be drawn if the collection is large, and contains many species (a practical limit is approximately \(S > 30\)). Basically, a species-abundance distribution (figure 3A) plots the number of species that are represented by \(r = 0, 1, 2, \ldots\) individuals against the abundance \(r\). Thus, in figure 3A there were 25 species with 1 individual, 26 species with 2 individuals, etc. More often than not,
the species are grouped in logarithmic density classes. Thus one records the number of species with density e.g. between \(1\) and \(e^{0.5}\), between \(e^{0.5}\) and \(e^{1}\), etc. (figure 3B). A practice, dating back to PRESTON (1948), is to use logarithms

![Figure 1: Ranked species abundance curves, representing the same data, with none, one or both axes on a logarithmic scale.](image)
to the base 2. One then has the abundance boundaries 1, 2, 4, 8, 16, etc. These so-called “octaves” have two disadvantages. The class boundaries are integers, which necessitates decisions as to which class a species with an abundance

Figure 2: The same data set represented as a k-dominance curves (A-B), the Lorenzen curve (C) and the collector's curve (D).

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equal to a class mark belongs; and, the theoretical formulation of models is "cluttered" (MAY, 1975) by factors ln(2), which would vanish if natural logs were used.

Figure 3: The same data set represented as species abundance distributions (A-C), and cumulative species abundance on a probit scale (D).
The ordinate of species-abundance distributions may be linear or logarithmic. Often one plots the cumulative number of species in a density group and all less abundant density groups on a probit scale (figure 3D).

2.2. Species-abundance models

Two kinds of models have been devised to describe the relative abundances of species. "Resource apportioning models" make assumptions about the division of some limiting resource among species. From these assumptions a ranked abundance list or a species-abundance distribution is derived. The resource apportioning models have mainly historical interest. In fact, observed species-abundance patterns cannot be used to validate or discard a particular model, as has been extensively argued by Pielou (1975, 1981). One should consult these important publications before trying to validate or refute a certain model fortuitously!

"Statistical models" make assumptions about the probability distributions of the numbers in the several species within the community, and derive species-abundance distributions from these.

2.2.1. The niche preemption model (Geometric series ranked abundance list)

This resource apportioning model was originally proposed by Motomura (1932). It assumes that a species preempts a fraction \( k \) of a limiting resource, a second species the same fraction \( k \) of the remainder, and so on. If the abundances of the species are proportional to their share of the resource, the ranked-abundance list is given by a geometric series:

\[
\begin{align*}
  k, k(1-k), \ldots, k(1-k)^{(S-2)}, k(1-k)^{(S-1)}
\end{align*}
\]

where \( S \) is the number of species in the community. May (1975) derives the species abundance distribution from this ranked abundance list (see also Pielou, 1975).

The geometric series yields a straight line on a plot of log abundance against rank. The communities described by it are very uneven, with high dominance of the most abundant species. It is not very often found in nature. Whittaker (1972) found it in plant communities in harsh environments or early successional stages.

2.2.2. The negative exponential distribution (broken-stick model)

A negative exponential species abundance distribution is given by the probability density function:

\[
\psi(y) = Se^{-Sy}
\]

Stated as such, it is a statistical model, an assumption about the probability distribution of the numbers in each species. However, it can be shown (Webb,
that this probability density function can be arrived at via the "broken-stick model" (MacArthur, 1957). In this model a limiting resource is compared with a stick, broken in S parts at S - 1 randomly located points. The length of the parts is taken as representative for the density of the S species subdividing the limiting resource. If the S species are ranked according to abundance, the expected abundance of species i, N_i is given by:

\[ E(N_i) = \frac{1}{S} \sum_{x=i}^{S} \frac{1}{x} \]  

(2.5)

The negative exponential distribution is not often found in nature. It describes a too even distribution of individuals over species to be a good representation of natural communities. According to Frontier (1985) it is mainly appropriate to describe the right-hand side of the rank frequency curve, i.e. the distribution of the rare species. As these are the most poorly sampled, their frequencies depend more on the random elements of the sampling than on an intrinsic distribution of the frequencies.

Pielou (1975, 1981) showed that a fit of the negative exponential distribution to a field sample does not prove that the mechanism modelled by the broken-stick model governs the species-abundance pattern in the community. Moreover, the broken-stick model is not the only mechanism leading to this distribution. The same prediction of relative abundance can be derived by at least three other models besides the niche partitioning one originally used (Cohen, 1968; Webb, 1974).

The observation of this distribution does indicate (May, 1975) that some major factor is being roughly evenly apportioned among the community's constituent species (in contrast to the lognormal distribution, which suggests the interplay of many independent factors).

2.2.3. The log-series distribution

The log-series was originally proposed by Fisher et al. (1943) to describe species abundance distributions in large moth collections. The expected number of species with r individuals, E_r, is given as:

\[ E_r = \alpha \frac{X^r}{r} \]  

(2.6)

(r = 1, 2, 3, ...). \( \alpha (> 0) \) is a parameter independent of the sample size (provided a representative sample is taken), for which X (0 < X < 1) is the representative parameter. The parameters \( \alpha \) and X can be estimated
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by maximum likelihood (Kempton & Taylor, 1974), but are conveniently estimated as the solutions of:

\[ S = -\alpha \ln(1 - x) \]  

and

\[ N = \frac{\alpha X}{1 - X} \]  

The parameter \( \alpha \), being independent of sample size, has the attractive property that it may be used as a diversity statistic (see further). An estimator of the variance of \( \alpha \) is given as (Anscombe, 1950):

\[ \text{var}(\alpha) = \frac{\alpha}{\ln X(1 - X)} \]  

Kempton & Taylor (1974) give a detailed derivation of the log-series distribution. It was fitted to data from a large variety of communities (e.g. Williams, 1964; Kempton & Taylor, 1974). It seems, however, to be in general less flexible than the log-normal distribution. In particular, it cannot account for a mode in the species-abundance distribution, a feature often found in a collection. According to the log-series model, there are always more species represented by 1 individual than there are with 2. The truncated log-normal distribution can be fitted to samples with or without a mode in the distribution.

Caswell (1976) derived the log-series distribution as the result of a neutral model, i.e. a model in which the species abundances are governed entirely by stochastic immigration, emigration, birth and death processes, and not by competition, predation or other specific biotic interactions. He proposes to use this distribution as a “yardstick”, with which to measure the occurrence and importance of interspecific interactions in an actual community. Other models have been proposed to generate the log-series distribution (Boswell & Patil, 1971) but they all contain the essentially neutral element as to the biological interactions. However, the proof that any form of biological interaction will yield deviation from the log-series is not given. Neither is it proven that “neutral” communities cannot deviate from the log-series. Therefore we think that the fit of this distribution cannot be considered as a waterproof test for species interactions.

2.2.4. The log-normal distribution

Preston (1948) first suggested to use a log-normal distribution for the description of species-abundance distributions. It was shown by May (1975) that a log-normal distribution may be expected, when a large number of independent environmental factors act multiplicatively on the abundances of the species (see also Pielou, 1975).
When the species-abundance distribution is log-normal, the probability density function of \( y \), the abundance of the species, is given by:

\[
\psi(y) = \frac{1}{y\sqrt{2\pi V_z}} \exp \left(-\frac{(\ln y - \mu_z)^2}{2V_z}\right)
\]  \hspace{1cm} (2.10)

The mean and variance of \( y \) are:

\[
\mu_y = \exp \left(\mu_z + \frac{V_z}{2}\right) \hspace{1cm} (2.11)
\]
\[
V_y = (\exp(V_z) - 1) \exp(2\mu_z + V_z) \hspace{1cm} (2.12)
\]

where \( \mu_z \) and \( V_z \) are the mean and variance of \( z = \ln(y) \).

If the species abundances are lognormally distributed, and if the community is so exhaustively sampled that all the species in the community (denoted \( S^* \)) are represented in the sample, the graph of the cumulative number of species on a probit scale (figure 3D) against log abundance will be a straight line. This is not normally the case.

In a limited sampling a certain number of species \( S^* - S \) will be unrepresented in the sample (\( S \) being the number of species in the sample). The log-normal distribution is said to be truncated. In the terminology of PRESTON (1948) certain species are hidden behind a "veil line". It follows that it is not good practice to estimate the parameters of the lognormal distribution from a cumulative plot on a probit scale. In fact if one does not estimate the number of unsampled species, it is impossible to estimate the proportion of the total number of species in a particular log density class. Species abundances that are lognormally distributed will not yield straight lines if one takes into account only the species sampled. Note also that the normal regression analysis is not applicable to highly correlated values such as cumulative frequencies. (If the frequencies are replaced by evenly distributed random numbers, their cumulative values on probit scale still yield very "significant" correlations with log abundance).

In fitting the log-normal two procedures are used (apart from the wrong one already discussed). The conceptually most sound method is to regard the observed abundances of species \( j \) as a Poisson variate with mean \( \lambda_j \), where the \( \lambda_j \)'s are lognormally distributed. The probability, \( p_r \), that a species contains \( r \) individuals is then given by the Poisson log-normal distribution (see BULMER, 1974). \( p_r \) can be solved approximately for \( r > 10 \), but must be integrated numerically for smaller values of \( r \). BULMER (1974) discusses the fitting to the data by maximum likelihood. PIELOU (1975) argues that the fitting of the Poisson lognormal, though computationally troublesome, is not materially better than the alternative procedure, consisting in the direct fitting of the
continuous lognormal. The complete procedure in recipe-form is given in Pielou (1975).

2.3. On fitting species-abundance distributions

Ever since Fisher et al. (1943) used the log-series, and Preston (1948) proposed the log-normal to describe species-abundance patterns, ecologists have been debating which model is the most appropriate. Especially the log-normal and the log-series have (had) their fan-clubs (e.g. Shaw et al., 1983; Gray, 1983 and other papers). In our opinion, these debates are spurious. As Pielou (1975) remarked, the fact that e.g. the log-normal fits well in many instances, tells us more about the versatility of the log-normal than about the ecology of these communities. Although most of the distributions have a kind of biological rationale (to make them more appealing to a biological audience?) the fact that they fit does not prove that the “biological” model behind them is valid in the community. The fitting of a model to field data is meaningful if the parameter estimates are to be used in further analysis. This is analogous to the use of the normal distribution in ANOVA: in order to perform an ANOVA, the data should be normally distributed. Of course this must be checked, but only as a preliminary condition. No one draws conclusions from the fit or non-fit of the normal distribution to experimental data, but from the test performed afterwards. Similarly, if a particular model fits reasonably well to a set of field data, the parameter estimates can be used, e.g. in respect to the diversity of the communities.

3. Diversity indices derived from species-abundance distributions

Historically, the first diversity measure was derived by Fisher et al. (1943) as a result of the derivation of the log-series distribution. The parameter $\alpha$ of the log-series distribution is independent of sample size. From equation (2.7) it is easily seen that $\alpha$ describes the way in which the individuals are divided among the species, which is a measure of diversity. In adopting the log-series model for the species-abundance distribution, the evenness is already specified, so that $\alpha$ only measures the relative species richness of the community. $\alpha$, as determined by the fitting of the log-series model to the sample, is only valid as a diversity index when the log-series fits the data well. The same reasoning can be extended to the log-normal distribution. Preston (1948) expressed the diversity (richness) as the (calculated) total number of species in the community, $S^*$. 

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The use of the log-series $\alpha$ was taken up again, and extended by Kempton & Taylor (1974). Taylor et al. (1976) showed that, when the log-series fits the data reasonably well, $\alpha$ has a number of attractive properties. The most important of these are that (compared to the information statistic $H'$ and Simpson's index; see below) it provided a better discrimination between sites, it remained more constant within each site (all sites were sampled in several consecutive years), it was less sensitive to density fluctuations in the commonest species, and it was normally distributed. On the other hand, when the data deviate from the log-series, $\alpha$ is more dependent on sample size than the other indices.

4. Rarefaction

An obvious index of species richness is the number of species in the sample. However, it is clear that this measure is highly correlated with sample size, an undesirable property. Sanders (1968) proposed a method to reduce samples of different sizes to a standard size, so as to make them comparable in terms of the number of species. The formula used by Sanders (1968) was corrected by Hurlbert (1971), who showed that the expected number of species in a sample of size $n$ is given by:

$$ES_n = \sum_{i=1}^{S} \left( 1 - \frac{N - N_i}{n} \right)$$

where $N_i$ is the number of individuals of the $i$-th species in the full sample, which had sample size $n$ and contained $S$ species. The notation in square brackets $\binom{A}{B}$ indicates the number of permutations of $A$ elements in groups of size $B$. Alternatively, random samples can be drawn by computer from the original sample (Simberloff, 1972). For an example of application of this method to deep-sea benthos see Soetaert & Heip (1990).

5. Hill's (1973) diversity numbers

Hill (1973) provided a generalized notation that includes, as a special case, two often used heterogeneity indices. Hill defined a set of "diversity numbers"
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of different order. The diversity number of order $a$ is defined as:

$$H_a = \left( \sum_i p_i^a \right)^{1/(1-a)}$$  \hspace{1cm} (5.1)

where $p_i$ is the proportional abundance of species $i$ in the sample. In the original notation $N$ is used instead of $H$, but to avoid confusion with abundance $N$, we propose to use $H$ (Hill) instead. For $a = 0$, $H_0$ can be seen to equal $S$, the number of species in the sample. For $a = 1$, $H_1$ is undefined by equation (5.1). However, defining

$$H_1 = \lim_{a \to 1} H_a$$ \hspace{1cm} (5.2)

it can be shown that

$$H_1 = \exp(H')$$ \hspace{1cm} (5.3)

where $H'$ is the well-known Shannon-Wiener diversity index:

$$H' = -\sum p_i \ln p_i$$ \hspace{1cm} (5.4)

This is the most widely used diversity index in the ecological literature.

Note that in the usual definition of the Shannon-Wiener diversity index logarithms to the base 2 are used. Diversity then has the peculiar units "bits·ind$^{-1}$".

The diversity number $H_1$ is expressed in much more natural units. It gives an equivalent number of species, i.e. the number of species $S'$ that yields $H_1$ if all species contain the same number of individuals, and thus if all $p_i = 1/S'$. This can be seen in equation (5.3), which in this case reverts to:

$$H_1 = \exp \left( -\ln(1/S') \right) = S'$$ \hspace{1cm} (5.5)

An additional advantage of $H_1$ over $H'$ is that it is approximately normally distributed.

It has been argued (see e.g. Pielou, 1975) that for small, fully censused communities the Brillouin index should be used. This index is given by:

$$H = \frac{1}{N} \log \frac{N!}{\prod N_i!}$$ \hspace{1cm} (5.6)

in which $\prod N_i = N_1 \cdot N_2 \cdots N_S$.

We do not recommend this index. The theoretical information argument for its use should be regarded as allegoric: it has no real bearing to ecological theory. Peet (1974) showed with an example that the Brillouin index has counter-intuitive properties: depending on sample size, it can yield higher values for less evenly distributed communities.
The next diversity number, $N_2$, is the reciprocal of Simpson's dominance index $\lambda$, which is given by:

$$\lambda = \sum p_i^2$$

(5.7)

for large, sampled, communities. If one samples at random and without replacement 2 individuals from the community, Simpson's index expresses the probability that they belong to the same species. Obviously, the less diverse the community is, the higher is this probability. In small, fully censused communities, the correct expression for Simpson's index is:

$$\lambda = \sum \frac{N_i(N_i - 1)}{N(N - 1)}$$

(5.8)

where $N_i = \text{number of individuals in species } i$, $N$ is the total number of individuals in the community.

In order to convert Simpson's dominance index to a diversity statistic it is better to take reciprocal $1/\lambda$, as is done in Hill's $H_2$, than to take $1 - \lambda$. In that way the diversity number $H_2$ is again expressed as an equivalent number of species.

Hill (1973) pointed out that $\lambda$ is a weighted mean proportional abundance, as it can be written as:

$$\lambda = \frac{\sum w_i p_i}{\sum w_i}$$

(5.9)

where the weights are equal to the relative abundance $w_i = p_i$.

The diversity number of order $+\infty$, $H_{+\infty}$, is equal to the reciprocal of the proportional abundance of the commonest species. It is also called the "dominance index". May (1975) showed that it characterises the species-abundance distribution "as good as any, and better than most" single diversity indices. It is also the most easily estimated diversity number since its calculation only requires distinction between the commonest species and all the others.

Hill (1973) showed that the diversity numbers of different orders probe different aspects of the community. The number of order $+\infty$ only takes into account the commonest species. At the other extreme, $H_{-\infty}$ is the reciprocal of the proportional abundance of the rarest species, ignoring the more common ones. The numbers $H_0$, $H_1$, and $H_2$ are in between in this spectrum. $H_2$ gives more weight to the abundance of common species (and is, thus, less influenced by the addition or deletion of some rare species) than $H_1$. This, in turn, gives less weight to the rare species than $H_0$, which, in fact, weighs all species equally, independent of their abundance. It is good practice to give diversity numbers of different order when characterising a community. Moreover, these numbers are useful in calculating evenness (see below).
6. The subdivision of diversity

6.1. Hierarchical subdivision

In the calculation of diversity indices, all species are considered as different, but equivalent: one is not concerned with the relative differences between species. However, in nature some species are much more closely related to some other species than to the rest of the community. This relation may be considered according to different criteria, e.g. taxonomic relationships, general morphological types, trophic types, etc. It may therefore be desirable to subdivide the total diversity in a community in a hierarchical way. Pielou (1969) shows how the Shannon-Wiener diversity $H'$ can be subdivided in a hierarchical way. The species are grouped in genera, and the total diversity equals:

$$H'_T = H'_g + H'_{wg}$$  \hspace{1cm} (6.1)

where $H'_g$ is the between genera diversity given by:

$$H'_g = - \sum_i q_i \log q_i$$  \hspace{1cm} (6.2)

and

$$H'_{wg} = \sum_i q_i \left( - \sum_j r_{ij} \log r_{ij} \right)$$  \hspace{1cm} (6.3)

is the average within-genus diversity. The same procedure may be repeated to partition the between-genera diversity into between-families and average within-family diversity. This approach was generalised by Routledge (1979) who showed that the only diversity indices that can be consistently subdivided are the diversity numbers of Hill (1973) (of which $H'$ can be considered a member, taking into account the exponential transformation).

The decomposition formula is:

$$\left( \sum_i \sum_j r_{ij}^{a} \right)^{1/(1-a)} = \left( \sum_i q_i^{a} \right)^{1/(1-a)} \left( \sum_i q_i^{a} \sum_j r_{ij}^{a} \right) / \sum_i q_i^{a}$$  \hspace{1cm} (6.4)

for $a \neq 1$.

In equation (6.3) $q_i = $ proportional abundance of group (e.g. genus) $i$, $r_{ij} = $ proportional abundance of species $j$ in group $i$, $t_{ij} = $ proportional abundance of species $j$ (belonging to group $i$) relative to the whole community.

It can be seen that the community diversity is calculated as the product of the group diversity and the average diversity within groups, weighted by the
proportional abundance of the groups. Note that this is consistent with Pielou’s formulae (eq. (5.9)) since $H_1 = \exp(H')$.

The hierarchical subdivision of diversity may be useful to study the differences in diversity between two assemblages, and to investigate whether a higher diversity in one assemblage can be attributed mainly to the addition of some higher taxa (suggestive of the addition of new types of niches), or of a diversification of the same higher taxa that are present in the low-diversity assemblage.

It may also be useful to study other than taxonomic groups. Natural ecological groupings, such as the feeding or body types may be particularly interesting. Heip et al. (1984, 1985) used $\theta$ as a “trophic diversity index” to describe the diversity in feeding types of nematodes,

$$\theta = \sum_{i=1}^{n} q_i^2$$

(6.5)

where $q_i$ is the proportion of feeding type $i$ in the assemblage and $n$ is the number of feeding types.

6.2. Spatio-temporal diversity components

All ecological communities are variable at a range of spatio-temporal scales. Thus if one examines a set of samples, (necessarily) taken at different points in space, and possibly also in time, and calculates an overall diversity index, it is unclear what is actually measured. Whereas diversity may be small in small patches at a particular instant, additional diversity may be added by the inclusion in the samples of diversity components due to spatial or temporal patterns.

Following Whittaker (1972) one often distinguishes between $\alpha$-diversity, the diversity within a uniform habitat (patch), $\beta$-diversity, the rate and extent of change in species composition from one habitat to another (e.g. along a gradient), and $\gamma$-diversity, the diversity in a geographical area (e.g. the intertidal range of a salt marsh). These are useful and important distinctions.

The subdivision of total diversity $H'$ in ecological components is discussed by Allen (1975). He treats a sampling scheme where $S$ species are sampled in $q$ sites, each consisting of $r$ microhabitats. The problem is different from a hierarchical subdivision, since the same species may occur in different microhabitats and sites (it can, of course, only belong to one genus, one family, etc. in hierarchical subdivision). Allen (1975) presents two solutions. One can treat the populations of the same species in different microhabitats as the fundamental entities. Total diversity is then calculated on the basis of the
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proportional abundance (in relation to the total abundance in the study) of these populations. This total diversity can then be subdivided hierarchically.

Alternatively, one can subdivide the species diversity in the total study in average within microhabitat diversity, average between microhabitat (within site) diversity, and average between site diversity components. The latter computations are generalised for Hill’s (1973) diversity numbers by Routledge (1979).

6.3. Cardinal and ordinal diversity measures

Species are different and fulfil different roles in ecosystems, and within species individuals are different as well. Since most diversity indices are based on the relative abundance of the different species representing the community, abundance is the only trait of species that is considered to differ between them. Cousins (1991) distinguishes between indices that treat each species as equal (cardinal indices) and those that treat each species as essentially different.

Species that are taxonomically more similar are also more similar in their morphology, and often in their behaviour and their ecological role in the system, than species that belong to different higher taxa. In practice diversity indices are often applied only to certain taxonomic groups (taxocenes) and the precise taxon level depends on the group being studied. Vane-Wright et al. (1991, see also May, 1990) have explored the implications of measures of taxonomic distinctiveness. They have used the hierarchical taxonomic classification to calculate an “information index” for species that is based on the number of branch points in the classification tree.

The idea has been taken one step further by Warwick & Clarke (1995) who introduced two new indices. In the first one, called taxonomic diversity, the abundance of a species is weighed with the taxonomic path length linking the species with the other species. Taxonomic diversity is the average (weighted) path length in the taxonomic tree between every pair of individuals. A second index, called taxonomic distinctness, is defined as the ratio between the observed taxonomic diversity and the value that would be obtained if all individuals belong to the same genus. This index was shown to be very sensitive to changes in community composition of macrobenthos around drilling platforms in the North Sea.

When diversity is represented by ranking each species in an order of some kind, the resulting index is called an ordinal index by Cousins (1991). The classical indices, such as Shannon-Wiener and Simpson’s index, are cardinal, whereas species abundance distributions, size spectra and species lists are ordinal representations. Cardinal indices are proposed to be useful for describing the diversity of a guild of species or the species within certain classes of body
size or weight, but are considered unsuitable for description of entire communities, where ranking the species is the better option.

7. Sampling properties of diversity indices

Since estimates of the true population (in a statistical sense) value are based on sampling that population it is necessary to pose the question what the sampling properties of diversity indices are. The sampling method itself has to fulfill a number of conditions and usually requires randomness. A good estimator of an index must be unbiased with minimum variance. As already pointed out a number of indices are biased, e.g. all indices based on estimates of S, the number of species in the community. The estimator of the Shannon-Wiener index is also biased. Estimators of the Simpson index and the rarefaction measure on the contrary appear to be unbiased. Species must be distributed at random and independent of other species. This is not usually the case and there is as yet no method that will produce unbiased diversity indices with low sampling variance and sampling distributions not influenced by species distribution patterns in the field.

Two methods that have become increasingly popular to overcome some of these difficulties are the jackknife and the bootstrap methods. These are resampling methods and are discussed in more detail by Dallot (this issue).

In the jackknife method pseudovalues are computed for the parameter of interest (e.g. species number, or Hill H1) which measure the weighted influence of each sample. The i-th pseudo-value is

\[ g_i = n g_0 - (n - 1) g_{-i} \]  

(7.1)

in which \( g_0 \) is the parameter computed with the \( n \) samples pooled and \( g_{-i} \) the corresponding value omitting the i-th sample.

In the bootstrap method the parameter is computed using a set of observed values drawn with replacement from the original set.

8. Evenness

The distribution of individuals over species is called evenness. It makes sense to consider species richness and species evenness as two independent characteristics of biological communities that together constitute its diversity (Heip, 1974).
Several equations have been proposed to calculate evenness from diversity measures. The most frequently used measures, which converge for large samples (Peet, 1974) are:

\[ E = \frac{I - I_{\text{min}}}{I_{\text{max}} - I_{\text{min}}} \]  

(8.1)

and

\[ E = \frac{1}{I_{\text{max}}} \]  

(8.2)

where \( I \) is a diversity index, and \( I_{\text{min}} \) and \( I_{\text{max}} \) are the lowest and highest values of this index for the given number of species and the sample size.

To this class belongs Pielou's \( J \):

\[ J = \frac{H'}{H'_{\text{max}}} = \frac{H'}{\log S} \]  

(8.3)

The condition of independence of evenness measures from richness measures is not fulfilled for the most frequently used evenness indices, such as \( J' \) (surprisingly, this is still the most widely used evenness index despite twenty years of literature describing its poor performance). As discussed by Peet (1974) such measures depend on a correct estimation of \( S^* \), the number of species in the community. It is quasi impossible to estimate this parameter. Substituting \( S \), the number of species in the sample, makes the evenness index highly dependent on sample size. It also becomes very sensitive to the near random inclusion or exclusion of rare species in the sample.

Hill (1973) proposed to use ratios of the form:

\[ E_{a:b} = \frac{N_a}{N_b} \]  

(8.4)

as evenness indices (where \( N_a \) and \( N_b \) are diversity numbers of order \( a \) and \( b \) respectively). Note that \( H' - H'_{\text{max}} = \ln(N_1/N_0) \) belongs to this class, but that \( J' = H'/H'_{\text{max}} \) does not. These ratios are shown to possess superior characteristics, compared with \( J' \). Hill (1973) also showed that in an idealised community, where the hypothesised number of species is infinite and the sampling is perfectly random, \( E_{1:0} \) is always dependent on sample size. \( E_{2:1} \) stabilises, with increasing sample size, to a true community value. However, in practice all measures depend on sample size.

\[ E_{1:0} = \frac{e^{H}}{S} \]  

(8.5)

Heip (1974) proposed to change the index to

\[ E'_{1:0} = \frac{e^{H} - 1}{S - 1} \]  

(8.6)
In this way the index tends to 0 as the evenness decreases in species-poor communities. Due to a generally observed correlation between evenness and number of species in a sample, $E_{1:0}$ tends to 1 as both $e^H \rightarrow 1$ and $S \rightarrow 1$. However this index falls into the same category as $J$, being dependent on an estimate of $S$.

A whole series of evenness indices can be derived from Simpson’s dominance index $\lambda$. Since the maximum value of $\lambda$ is $1/S$ ($S =$ number of species), an evenness index can be written as

$$E = \frac{1/\lambda}{S}$$

(8.7)

This corresponds to $E_{2:1}$ of Hill (1973)

$$E_{2:1} = \frac{1/\lambda}{e^H}$$

(8.8)

which was modified by Alatalo (1981) in the same way as Heip (1974) modified $E_{1:0}$.

$$E'_{2:1} = \frac{1/\lambda - 1}{e^H - 1}$$

(8.9)

Even in the recent literature (Smith & Wilson, 1996) it is recognized that the measurement of evenness is still very much a matter of debate and the literature continues to be “plagued” by new propositions (Molinaro, 1989; Camargo, 1992; Nee et al., 1992; Bulla, 1994). If the criterion of independence of measures for species richness and evenness (Heip, 1974) is accepted, the choice of indices becomes more restricted. A good discussion is given by Smith & Wilson (1996) who applied a series of additional requirements, e.g. that the index should decrease by reduction in abundance of minor species, decrease by addition of one very minor species, be unaffected by the units used, etc. These authors concluded that the independence of richness criterion is the only sensible one and only five indices passed this test.

However, Smith & Wilson’s (1996) comparison is valid for samples only and several of the indices proposed are still dependent on the number of species $S$ in the community and therefore on sample size. Still, their idea to use the variation in species abundance is attractive (Hill, 1997). If one uses Hill’s number $H_2 = 1/\lambda$ a simple statistic is the weighted mean-square deviation from the proportional abundances that would be expected for $H_2$ equally abundant species. A measure of evenness is then:

$$D_{MS} = \frac{\sum w_i (p_i - \lambda)^2}{\sum w_i}$$

(8.10)
in which $\text{MS} = \text{mean square}$, $\lambda$ is Simpson's index (eq. (5.9)) and $w_i = p_i$ (eq. (2.2)).

Hill (1997) also shows that the expected mean and variance of the relative abundance $p_i$ are given by

$$\begin{align*}
E(p_i) &= \lambda \\
\text{Var}(p_i) &= \text{DMS}
\end{align*}$$

(8.11)

(8.12)

A measure of the shape of the species abundance relation is given by

$$D_{\text{MS}}^* = \text{DMS}/\lambda^2$$

(8.13)

and a measure of evenness by:

$$E_{\text{MS}} = 1(1 + D_{\text{MS}}^*)$$

(8.14)

In general, species-abundance distributions show more information about the evenness than any single index. On the other hand, statistics describing these distributions can also be used as measures of evenness. Examples of indices that perform well are the one proposed by Camargo (1992), based on the variance in abundance over the species and the one proposed by Smith & Wilson (1996).

$$E_q = (-2/\pi) \arctan b'$$

(8.15)

in which $b'$ is the slope of the scaled rank of abundance on log abundance fitted by least square regression. The reader is referred to Smith & Wilson (1996) for further details.

9. The choice of an index

The choice of an index has to be considered with care. In our opinion Hill's diversity numbers present a coherent system for diversity estimates. They provide numbers that are equivalent to species numbers and include the simplest measure of species richness, the number of species in the sample as a special case. They also include variants of the Shannon-Wiener and the Simpson indices to which most use of diversity indices has converged. These indices reflect both the evenness (as they are based on the relative abundance of the species considered $p_i$) and species richness (as they sum up over all the species in the sample). They have even been called evenness indices in the recent literature (Wilsey & Potvin, 2000), an idea that is worth exploring but very much in contrast to established use of the term.

Lamshead et al. (1983) have noted that, whenever two $k$-dominance curves do not intersect all diversity indices will yield a higher diversity for the
sample represented by the lower curve. In such a case one could even try using Hill’s diversity number $+\infty$ (the relative abundance of the most dominant species), for instance in monitoring or impact studies where the need for “quick and dirty” measures is often required for reasons of cost. Equivocal results arise as soon as the $k$-dominance curves intersect.

The different measures of diversity are more sensitive to either the commonest or the rarest species. An elegant approach to the analysis of this sensitivity is provided by the response curves of Peet (1974). In order to summarise the diversity characteristics of a sampled community, it is advisable to provide the diversity numbers $N_0, N_1, N_2$, possibly also $N_{+\infty}$, the dominance index. If permitted by the sampling scheme, one can use these indices in a study of hierarchical and/or spatio-temporal components of diversity. In any case, it should be remembered that the indices depend on sample size, sample strategy (e.g. location of the samples in space and time), spatio-temporal structure of the community, and sampling error. Although formulae for the estimation of the variance of $H'$ have been proposed, these do not include all these sources of error (Heip & Engels, 1974; Frontier, 1985).

Evenness indices should still be regarded with caution, but the latest propositions by Smith & Wilson (1996) and Hill (1997), although perhaps conflicting, deserve further study. It is always advisable to use species-abundance plots to study evenness.

Finally, we should stress the possibilities and limitations of diversity and evenness indices. An index must be regarded as a summary of a structural aspect of the assemblage. As has been stressed throughout this article, different indices summarise slightly different aspects. In comparing different assemblages, it is useful to compare several indices: this will indicate specific structural differences. A diversity index summarises the structure, not the functioning of a community. It is thus very well possible that two assemblages have a similar diversity, whereas the mechanisms leading to their structures are completely different (e.g. Coull & Fleeger, 1977). Often these functional aspects cannot readily be studied by observing resultant structures, and may require an experimental approach.

References


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