A Random Categorization Model for Hierarchical Taxonomies

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A taxonomy is a standardized framework to classify and organize items into categories. Hierarchical taxonomies are ubiquitous, ranging from the classification of organisms to the file system on a computer. Characterizing the typical distribution of items within taxonomic categories is an important question with applications in many disciplines. Ecologists have long sought to account for the patterns observed in species-abundance distributions (the number of individuals per species found in some sample), and computer scientists study the distribution of files per directory. Is there a universal statistical distribution describing how many items are typically found in each category in large taxonomies? Here, we analyze a wide array of large, real-world datasets – including items lost and found on the New York City transit system, library books, and a bacterial microbiome – and discover such an underlying commonality. A simple, non-parametric branching model that randomly categorizes items and takes as input only the total number of items and the total number of categories is quite successful in reproducing the observed abundance distributions. This result may shed light on patterns in species-abundance distributions long observed in ecology. The model also predicts the number of taxonomic categories that remain unrepresented in a finite sample.

Hierarchical taxonomies are tree-like structures that organize items into subcategories, each of which descends along a unique path from a root category containing all items (Fig. 1(a)). From Aristotle to the Linnean classification of species, taxonomies have long been part of our way of classifying organisms, and more generally large corpora of knowledge.

We study the distribution of items among categories in a variety of large hierarchical classification systems, including disease incidence in 250 million patients1, bacterial microbiomes2, items for sale on www.amazon.com, books in the Harvard University library system, files per directory on a laptop, and items lost and found on the New York City transit system (Fig. 1(c) and Table 1). In all these datasets, a few categories are very popular while many categories contain only a few items. When categories are sorted by number of items and binned into a histogram, one can observe a range of shapes (Fig. 1(b)). As a function of the logarithm of the number of items, many datasets present a characteristic Gaussian or bell-shaped form – for patients per medical condition and www.amazon.com’s categorization – while others tend towards a highly skewed distribution with a long tail corresponding to many categories with low number of items – microbiome, files per directory, and the Harvard library books. This variation is correlated to how well-sampled the distribution is: datasets with a sufficiently large number of items relative to the number of categories are close to Gaussian, while those with many categories containing only a few items are skewed (Fig. 1(b)).

Results

Canonical lognormal distribution. The archetypal taxonomy is the categorization of organisms into species. Characterizing the distribution of the number of individuals of each species found in some area or sample – the so-called species abundance distribution (SAD) – is a long standing problem in ecology3. A classic observation is that SADs follow a “canonical lognormal distribution”, a lognormal distribution with standard deviation \( \sigma \) determined by the number of species \( k \):

\[
k = \frac{\sigma}{\ln 2} \sqrt{\frac{\pi}{2}} \exp(\sigma^2/2).
\]
However, in many real-world examples neither the canonical lognormal nor a lognormal with mean and variance chosen to best fit the data describe real-world SADs particularly well\textsuperscript{6,7}. In Fig. 2 we present such a dataset: a microbiome (a sample of a bacterial population categorized using Operational Taxonomic Units) for which the best-fit lognormal is a poor fit. In ecology, Sugihara\textsuperscript{5} proposed a “sequential broken stick” model to explain the origin of Eq. 1. The model randomly and sequentially divides items into subcategories, producing a branching tree. Sugihara’s approach to SAD was motivated by ecological niches and evolution, and he argued against a purely statistical explanation. However, if any set of items originally in one category are repeatedly and randomly subdivided, by the central limit theorem the logarithm of the number in any given subcategory should approach a normal distribution in the limit of many divisions, with a variance that depends on the number of divisions in a way closely related to Eq. 1. For this reason Sugihara’s model produces an approximation to a lognormal distribution, but the same would hold for any random process involving repeated subdivisions\textsuperscript{8}, including the one studied here.

As we will see, sequential random categorization appears to successfully account for the item abundance distribution in large hierarchical taxonomies, including many that have nothing to do with ecology. Random categorization can account for both the lognormal distributions observed in well-sampled datasets and the skewed shape of more poorly sampled taxonomies. In addition it can predict the detailed shape of the distribution (including its mean, variance, and the number of unsampled, empty categories) based only on the number of items and categories, entirely independently of the nature of the items being categorized. Of course, there are many models of random categorization in addition to the one studied in\textsuperscript{5}. Here, we focus on what we regard as the simplest such model. It is distinguished by the fact that within a class of related models, it produces the item
distribution with the minimum possible variance (see “More general random subdivision models” for details). The only inputs to the model are the total number of categories $k$ and total number of items $n$ – it contains no continuous or adjustable parameters. For the reason discussed above, when $n$ is sufficiently large compared to $k$ the distribution approaches lognormal with a mean and variance that depend on $k$ and $n$ (Fig. 3). We compare this model to four standard distributions characterized by one or two continuous parameters that we fit to the data: lognormal, geometric, log series, and negative binomial. The random categorization model is quite successful in reproducing our data compared to these distributions despite its lack of adjustable parameters (see “Comparison with data” for details).

**Random categorization model.** The model is defined as follows. Beginning with a single root category, it divides it into two subcategories, then randomly selects one subcategory and divides it into two, etc., producing...
a random binary tree (Fig. 1(a)). After \( q - 1 \) divisions the tree has \( q \) "leaves", each of which represents a category. Probabilities are then assigned to the leaves in a way that depends in a simple way on the number of branchings separating the leaf from the root. Finally, we "sample" the tree by populating the leaves with \( n \) items, distributed randomly according to the leaf probabilities.

The motivation for this last step is as follows. We regard the tree as representing a kind of “ideal” classification, while any real dataset is a sampling of that categorization. An example is the Library of Congress (LOC) categorization system, with \( q \) total categories. The collection of any individual library is a sampling of the full LOC collection. In most real libraries the sampling is sparse and only \( k < q \) of the LOC categories are represented by books in that library’s collection. If the number of items \( n \) is sufficiently large so that there are few or no empty categories, the model predicts an item abundance distribution that is indeed approximately lognormal (for reasons related to the central limit theorem and the statistics of the tree distribution). For smaller \( n \), the distribution of items in non-empty categories differs substantially from lognormal. In this way the same model – one with no continuous parameters and no fitting to the data – can produce distributions that look very different in different regimes, successfully reproducing the wide range represented in Fig. 3.

Figure 3. Histograms of the hierarchical taxonomic datasets. The plots are ordered by the value of the Shapiro-Wilk statistic W (measuring degree of normality). Grey is data, and we show kernel density estimates of the average of the random categorization model (blue curves) and the best fit lognormal (orange curves). In each plot the horizontal axis is the log of the fraction of items in the category, and the vertical axis is the log of the number of categories in each bin.
Generally, from the dataset we only know the number of non-empty categories $k$. To model the data, we therefore generate trees with $q \geq k$ categories. We choose $q$ so that the typical number of non-empty categories is $k$ when the tree is sampled with $n$ items (see Methods for details). For each dataset, the raw data – number of items per category – is sorted into a ranked list. The category with the largest number of items is ranked 1, while that with the least is ranked $k$. We then plot category rank versus the number of items (Figs 2, 4 and 5), or a binned histogram of the number of categories with some range of numbers of items (Fig. 3). The former is closely related to a graph of the cumulative distribution function of the item partitioning distribution, because for a category with rank $r$, $r/k$ is the probability that a random category would have as many or more items. The latter is related to the probability distribution function for the item distribution.

**Comparison with data.** The model can be thought of as producing a random partition of $n$ items into $k$. To compare the model to a particular dataset we use the model to generate many such partitions, and compute the relative entropy or Kullback-Leibler (KL) divergence of each to the data, generating a distribution of KL statistics (see “Comparison between theory and data”). The KL divergence is a measure of how informative the model is about the data – if it is small, it indicates that the model contains most of the information in the data. To compare
our model to a distribution characterized by continuous parameters, for each dataset we compute the KL statistics for partitions generated by \( k \) random draws from that distribution with parameters (mean and variance in the case of the lognormal distribution) fit to the data using maximum likelihood estimators (MLE). Because there are many categories with few items in the data, these fits are strongly affected by these categories (those with the highest rank). This explains why the fits are generally better for the least abundant categories (Fig. 4).

In six out of nine hierarchical datasets the random categorization model outperforms all four best-fit distributions, according to their median KL divergence. Of the three datasets for which one or more continuous distribution outperforms the random model, in one (CATH) the best model (lognormal) is very close to the random model. The other two (arxiv and NATC) have relatively small numbers of categories, which makes the variance in the KL divergence larger (Fig. 6 and Table 1). The success of the random model is quite remarkable given the model’s lack of adjustable parameters and suggests that the decisions made in designing hierarchical categorization systems follow a general pattern that our random categorization model is capturing.

As a test, we also compare the model to four categorizations that are not hierarchical taxonomies and where we would therefore not expect it to perform well: populations of German cities, U.S. counties, world countries, and the number of speakers per language. In contrast to the hierarchical datasets, the model is clearly inferior to the best-fit lognormal in all cases except for world population, where it is marginally inferior (Fig. 7).

An important point is that \( q \) can be thought of as a prediction for the total number of categories – that is, \( q - k \) is the number of categories that will not be represented in a typical sample of \( n \) items. This type of prediction has utility in a broad array of circumstances, and our model’s prediction exhibits a clear correlation to commonly employed estimators such as Chao9 (Table 1). An interesting test case is the Harvard library collection. The ratio of the total number of existing Library of Congress categories to \( k \) (the number of categories represented in

**Figure 5.** Rank-abundance plots for each dataset. Magenta points are data, light blue points are 100 realizations of the random model, whose average is the blue curve. Orange is the best fit lognormal, purple the best fit logseries, green the best fit geometric, and black the best fit negative binomial distribution. Plotted are the four non-hierarchical datasets (last four entries, Table 1).
Harvard’s collection) is approximately 16. This constitutes a lower bound on the true ratio, in the sense that the Library of Congress system may still be incomplete (certain subjects may remain as-yet unclassified, due to the dearth of extant books on those topics). Our model predicts a ratio of total categories to non-empty categories $q/k = 38$, while the Chao estimator predicts a ratio of 2.6. Hence, Chao dramatically under-predicts the number of empty categories. Our model may over-predict it, or it may correctly capture the current incomplete state of the system.

**Discussion**

Large hierarchical classification systems are ubiquitous, and our results illuminate a distinct and novel pattern in the distribution of items among categories in a broad array of such systems that cut across many fields. Randomly categorizing items in a binary tree, with no adjustable parameters and no input regarding the type of items being classified, appears to successfully reproduce this pattern. This result has broad applicability across many fields.
of science, and could serve as a starting point – a kind of null hypothesis – for more detailed investigations of item distributions. Deviations from the pattern observed here likely indicate that there is some significant effect that is not captured by random categorization: for instance preferential attachment\textsuperscript{10,11}, where added items are substantially more (or less) likely to be assigned to categories that are already large. The model can also be used to predict the number of empty or missing categories: given a finite sample of items, it makes a prediction for how many more categories there are than those represented in the sample. An important application of this prediction arises in ecology, where it can be used to estimate the number of rare species that were missed in some survey, and identifying the number of such “missing categories” clearly has applications far beyond ecology\textsuperscript{12}. Another feature of interest is that many of these datasets appear very close to power law over a large range of category ranks (Figs 2 and 4), and this behavior is correctly reproduced by the model. This may help account for the power laws previously observed in many contexts even without preferential attachment\textsuperscript{13}. A related discussion can be found in\textsuperscript{14}.

**Methods**

**Data description.** Our analysis makes use of nine datasets of hierarchical classifications (1–9) and four non-hierarchical datasets (10–14). From each dataset we extracted a list of categories and ranked it according to the number of items in each category (category 1 has the most items, category $k$ the fewest). Here we provide a brief description of each dataset and its source. A summary can be found in Table 1.
1. Harvard library: We counted the number of books per Library of Congress call number, extracted from the Harvard Library dataset published on http://library.harvard.edu/open-metadata#Harvard-Library-Bibliographic-Dataset. We considered call numbers that differed only after the “.” as equivalent.
2. Files: We counted the number of files in each directory on the Macbook Air laptop computer of the corresponding author, using the Bash command find -typed|whilereread -d|dir;ofind "$dir" -typef -maxdepth1 |wc -l|done >> |filedat. We checked that the file systems on several other Macintosh computers have a similar structure.
3. Microbiome: We used rank abundance of bacteria per operational taxonomic unit in a sample of the human oral microbiome, as described in http://jb.asm.org/content/192/19/5002.full#sec-142.
4. NATC: The number of books per category in the North American Title Count database, using a file sent to us by a librarian at the Library of Congress. The file is available upon request.
5. CATH: We took data from the file http://release.cathdb.info/latest_release/CathDomainList.v4.0.0, which is a database of protein structure classification. We define as a category the hierarchy of the first four levels of the classification, as described in http://www.ncbi.nlm.nih.gov/pmc/articles/PMC3525972/, and then count the number of domains in each category.
6. ArXiv: Using public APIs from the website www.arxiv.org, we counted the number of articles published in each subcategory in the year 2014. We considered only the primary category of each article, neglecting cross-listings.
7. MTA: We counted the number of items per category in the “lost and found” inventory of the Metropolitan Transportation Authority of New York City, as of December 24th, 2015. The data are published and constantly updated on http://advisory.mta.nyct.info/LPUWebServices/CurrentLostProperty.aspx, and we used the dataset downloaded and published on https://github.com/atmccann/mta-lost-found/blob/master/data/2014-12-24.json.
8. Amazon: We counted the number of items for sale on in departments with no subdepartments (that is, the lowest level categories) on www.amazon.com over a time period of several months in 2015–16.
9. Hospital patient medical conditions: We used a dataset of the number of patients per ICD-9 classified condition admitted to hospitals in the years 2000 to 2011. The source is the Observational Health Data Sciences and Informatics (OHDSI) dataset.
10. German cities: We used the population per each German city as of December 31st 2013, published by the Federal Statistical Office of Germany at https://www.destatis.de/DE/ZahlenFakten/LaenderRegionen/Regionales/Gemeindeverzeichnis/Administrativ/Archiv/GVAuszug/31122013_Auszug_GV.xls?__blob=publicationFile.
11. World population: We used the world population for 2014 published by the World Bank, at the website http://data.worldbank.org/indicator/SP.POP.TOTL.
12. US counties: We used the population for each county of the United States, as of the 2010 census, as published on “Annual Estimates of the Resident Population: April 1, 2010 to July 1, 2014, Source: U.S. Census Bureau, Population Division”.
13. Languages: We counted the number of native speakers for the languages of the world, published in the Encyclopedia of Language & Linguistics, at the page http://www.sciencedirect.com/science/article/pii/B0080448542090908.

Representing the distribution. To present the distribution of items among categories in these datasets we use several different graphical representations. One is the histogram of the number of categories with a given range of number of items (Fig. 3). This can be interpreted as the probability mass function (PMF) of the data – the probability that a randomly chosen category contains \( n \) items. This empirical PMF has two related shortcomings. First, its shape depends on the choice of binning. We use logarithmic binning, because the distribution is approximately lognormal for the well-sampled datasets. Second, because of sparse sampling, the tails of the distribution are noisy even with logarithmic binning.

Another representation of the data is a rank-frequency plot. Namely, we rank the categories by the number of items that fall into each (rank 1 being the category with the most items), and plot the rank as a function of the number of items (Figs 4 and 5). This graph is related to the cumulative distribution function (CDF) of the data: if a category has \( n \) items, its rank is proportional to the number of categories that have a number of items greater or equal to \( n \). For this reason the rank-frequency plot is an unbiased estimator of the CDF, and does not rely on a binning. Therefore, the tails of the distribution are accurately represented.

An alternative way to look at the data is to ask for the probability that a randomly chosen item will fall into a category with some given rank \( r \). The PMF of this probability is proportional to the quantile function, which is the functional inverse of the CDF – in other words, the rank-frequency plot with horizontal and vertical axes interchanged. This way of interpreting the data provides a robust estimator of a PMF, and we use it in comparing the model to the data.

Random categorization model. As described above, we introduce a random categorization model to explain the distribution of hierarchical taxonomies. The model has no continuous or adjustable parameters, and the only inputs are the number \( k \) of categories and the number \( n \) of items present in the dataset. It consists of the following steps:

1. Generate a random binary tree with \( q \geq k \) leaves;
2. Assign a probability to each leaf;
3. Randomly populate the leaves with \( n \) items according to these probabilities.
For step 1, we begin from a single root category that we divide into two child categories. Next, we choose one of these two categories at random and divide it into two, resulting in three categories. At each successive time step, we choose a category at random (with uniform probability) and divide it in two. We repeat this procedure until we reach the desired number of categories $q$ (“leaves” of the tree), which may be larger than $k$ (see below for details on how $q$ is determined).

To each leaf we assign a probability $p = 2^{-b+1}/X$, where $b$ is the depth of the leaf (the number of divisions separating that leaf from the root), $x$ is a uniform random variable between $-1$ and 1, and $X$ is a normalization factor that ensures the probabilities sum to one. The factor $2^{-b}$ corresponds to an equal expected division of the items in a parent category into each child category. Adding the random variable $x$ prevents the frequencies of items in well-sampled datasets from being artificially peaked at integer powers of two. We could also achieve a similar “smeared” distribution by assigning probabilities to each child category according to some continuous distribution rather than equally, or by dividing categories into a randomly varying number of child categories.

Of course, $b$ is itself a random variable that depends on the particular leaf and the structure of the random tree. Given a random tree with $q$ leaves, the probability that a randomly chosen leaf has depth $b$ is given by:

$$P(b, q) = 2^{-b} C(q - 1, b).$$

(2)

Here $C$ is an unsigned Stirling number of the first kind, which obeys the recursion relation:

$$C(n + 1, k) = nC(n, k) + C(n, k - 1)$$

with boundary conditions $C(n, 0) = 0$ for $n \geq 0$ and $C(n, k) = 0$ for $k \geq 0$, with $C(n, n) = 1$ for $n \geq 0$. From this and the definition of the Gamma function, we derive a recursion relation for the probability:

$$P(b, q + 1) = \frac{q - 1}{q + 1} P(b, q) + \frac{2}{q + 1} P(b - 1, q),$$

(4)

with boundary conditions $P(0, 1) = 1$, $P(0, q) = 0$ for $q \geq 0$. This recursion relation allows an efficient numerical calculation of the probabilities for arbitrary $b, q$, values.

This distribution is approximately normal in $b$ near its maximum, with variance $\sigma_b^2 = \sum_{j=2}^{q} (2j - 4j/(2j - 1)) \simeq -3.4 + 2 \ln q$ and mean $\mu_b = \sum_{j=2}^{q} 2j/(2j - 1) \simeq -0.85 + 2 \ln q$, where the approximations are valid at large $q$. Since the probabilities $p$ are related to $b$ by $p = 2^{-b}$, the variance in $\ln p$ is

$$\sigma_{\ln p} \simeq (\ln 2)^2 (3.4 + 2 \ln q).$$

(5)

Note that if $q$ is the number of species, the functional relation between the variance and the number of species is similar to the canonical lognormal Eq. 1 in that the logarithm of the number of species is linear in the variance at leading order, albeit with a different coefficient. This is a general feature of random tree models.

Having generated a tree and assigned probabilities, the last step is to assign $n$ items to the leaves by distributing them according to a multinomial distribution of $n$ over the $q$ probabilities $p_i$. There remains the question of determining the number of leaves $q$ that the tree should have. To do so, we notice that, given a tree with $q$ leaves, after multinomial sampling with $n$ items, we expect to get the following fraction of empty categories:

$$f_e(q, n) = \sum_{b=1}^{q} P(b, q) (1 - 2^{-b})^n,$$

(6)

with $P(b, q)$ given by eq. 2. If we wish to obtain an expected number $k$ of nonempty categories, we should choose $q$ by solving the nonlinear equation $k = q(1 - f_e(q, n))$. We use the value $q(k, n)$ that solves this equation for the number of categories we generate to model a dataset with given $k, n$.

These steps can be thought of as generating a random partition of $n$ into $q$. The probability distribution of these random partitions does not have a simple analytical expression as far as we are aware, but it is not difficult to sample numerically. To do so, we generate many random trees following the previous steps. For the datasets in which $q \gg k$, we only keep those trees where after populating with $n$ items, the number of nonempty categories is within 1% of $k$. Ideally we could keep only those with exactly the same number of non-empty categories as the data, but this would be time-consuming for large datasets and in any case does not affect the results significantly (see Fig. 8 for a robustness test comparing different values of this cutoff).

The code is implemented in Python, and it is available upon request from the authors. We have also made a web application available at the address https://rabadan.c2b2.columbia.edu/universaltaxdist in which the user can test the predictions of our model against any dataset of counts.

**More general random subdivision models.** There are many natural generalizations of this model. One is to assign unequal probabilities to the subcategories at each division. That is, for the binary model described above we assigned probabilities $p = 2^{-b+1}/X$ to the leaves, where $b$ is the number of branchings separating that leaf from the root and $x$ is a random variable ranging from $-1$ to 1. Apart from the smearing factor $x$, this corresponds to an equal ($50$–$50$) split at each binary subdivision. Instead, consider assigning probabilities to each of the two subcategories produced at a division according to some probability distribution $\rho(x)$ with $0 \leq x \leq 1$ (for instance, $\rho(x)$ could be a beta distribution), rather than equally. In this case, the variance of the log probability for a random leaf is the variance in the number of divisions $b$ times a factor of $\langle (\ln x)^2 \rangle = \int_0^1 (\ln x)^2 \rho(x) dx$. Note that
for the 50–50 model, $\langle (\ln x)^2 \rangle = (\ln \frac{1}{2})^2 = (\ln 2)^2$ (see Eq. 5). One can easily show that $(\ln 2)^2$ is the minimum possible value of $\langle (\ln x)^2 \rangle$ for distributions satisfying $\int_0^1 \rho(x)dx = 1$ and $\rho(x) = \rho(1 - x)$, as is necessary here. This shows that the model we have considered has the minimum possible variance within a large class of random division models.

Another generalization is to divide into $j$ subcategories according to some set of probabilities $p_j$, producing trees that are more general than the simple binary ($j = 2$ only) structure considered so far. For a model that always divides into a fixed number $j$ of subcategories (producing a “$j$-nary” tree rather than a binary tree), the mean and variance of $b$ (the number of branchings separating a random leaf from the root) for a tree with $q$ leaves are

$$\mu_j(q) = -\frac{j}{j-1} \left[ H\left(\frac{q}{j-1}\right) - H\left(\frac{j}{j-1}\right) \right],$$

where $H$ is a harmonic number, and

$$\sigma_j^2(q) = \frac{j}{(j-1)^2} \left[ (j-1) \left( \psi^{(0)}(1 + \frac{q}{j-1}) - \psi^{(0)}(2 + \frac{1}{j-1}) \right) + j \left( \psi^{(1)}(1 + \frac{q}{j-1}) - \psi^{(1)}(2 + \frac{1}{j-1}) \right) \right].$$

where $\psi^{(m)}(z)$ is a polygamma function of order $m$. If the probabilities are equally divided among the child categories so that $p = j^{-1}$, the mean and variance of the log probabilities are related to the above results simply by factors of $\ln j$ and $(\ln j)^2$ respectively (see Eq. 5). For $q \gg j$, the result is a variance in the log probabilities that is an increasing function of $j$. Hence, one sees that the minimum variance for a tree with a given number of leaves is achieved for the case of binary trees ($j = 2$).

A third generalization would be to choose to divide categories with a probabilities depending on the number of items in them (or the probability so far assigned to them), rather than uniformly. That is, one might consider a model where at each step large categories are more likely to be subdivided than small ones. This can produce an item distribution with lower variance than the model presented here, but we will not consider it further here. We have chosen to focus on the binary “equal divide” model because we regard it as the simplest, and it seems to do a surprisingly good job fitting the data.
Comparison between theory and data. We should think of the partition of items provided by the data as a single random sample of some underlying distribution. It is this underlying distribution that our model attempts to reproduce. In this situation it is difficult to use a null hypothesis frequentist test. A Bayesian inference approach would strongly depend on the chosen prior. It is possible to do a non-parametric test based on the empirical CDF, as for instance the Kolmogorov-Smirnov, Anderson-Darling or Mann-Whitney test. These are however sensitive to particular aspects of the frequency distribution, and would not properly express the degree by which our statistical model can capture the observed partition of items.

For these reasons, we characterize the degree of agreement of the data with our model (and the best-fit distributions we compare to) by calculating the Kullback-Leibler (KL) divergence, or relative entropy, between our model and the data. In general, given two probability distributions $p$ and $q$, the KL divergence is defined by

$$D_{KL}(p||q) = \int_{-\infty}^{+\infty} dx \ p(x) \ \log \frac{p(x)}{q(x)}$$

with a sum replacing the integral in the discrete case. This quantity is not symmetric in $p$ and $q$. Typically, $p$ represents a "fiducial" distribution, while $q$ is a model for or approximation of $p$. In the context of information theory, $D_{KL}(p||q)$ is the amount of information lost when the distribution $q$ is used to approximate $p$.

This approach to model comparison was used recently in ecology in the context of species-abundance distributions. There, $x$ is the rank of the species and $p$ and $q$ are estimates of the probability that a given individual belongs to the species with rank $x$ in the data and model respectively. Here, we employ a methodology closely related to that used by Alroy. To calculate $D_{KL}$ we estimate the probability that a random item belongs to a category with rank $x$ for both data and model. We estimate the probability for the data $p(x)$ and for each sampled tree $q(x)$ simply by the normalized fraction of items in the category with rank $x$. That is, $p(x) = n(x)/n$, where $n(x)$ is the number of items in the category of rank $x$ and $n$ is the total number of items, and the same for $q(x)$. We then compute $D_{KL}(p||q)$ using Eq. 9 and these estimates.

To compare the model to the data, we calculate the KL divergence for a set of 100 sampled trees, and then plot a histogram (actually, a kernel density estimate) of the results, showing the median KL divergence with a dashed vertical line. As a comparison, for each dataset we consider four alternative models: the lognormal, log series, negative binomial, and geometric distributions. In addition to the two integers $k$ and $n$ required to model the data, the lognormal and negative binomial distributions have two continuous parameters, while log series and geometric have one. The continuous parameters of these distributions are fit to the data $p(x)$ using maximum likelihood estimators. For each dataset we generate 100 sets of $k$ draws $q(x)$ from each of these distribution. Each set is then normalized so that $q(x)$ sums to 1, and we compute the KL divergence of each of these 100 sets relative to the data. The results are plotted in Figs 6 and 7.

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Author Contributions
G.D.A. performed the numerical analyses and collected most of the datasets. M.K. performed analytic calculations and wrote the manuscript, and is the corresponding author. R.R. provided the data that initiated the project and contributed to the figures.

Additional Information
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