Abstract
A new class of metrics appropriate for measuring effective similarity relations between sequences, say one type of similarity per metric, is studied. We propose a new "normalized information distance", based on the noncomputable notion of Kolmogorov complexity, and show that it minorizes every metric in the class (that is, it is universal in that it discovers all effective similarities). We demonstrate that it too is a metric and takes values in [0, 1]; hence it may be called the similarity metric. This is a theory foundation for a new general practical tool. We give two distinctive applications in widely divergent areas (the experiments by necessity use just computable approximations to the target notions). First, we computationally compare whole mitochondrial genomes and infer their evolutionary history. This results in a first completely automatic computed whole mitochondrial phylogeny tree. Secondly, we give fully automatically computed language tree of 52 different language based on translated versions of the "Universal Declaration of Human Rights".

1 Introduction
How do we measure similarity—for example to determine an evolutionary distance—between two sequences, such as internet documents, different language text corpora in the same language, among different languages based on example text corpora, computer programs, or chain letters? How do we detect plagiarism of student source code in assignments? The fast advance of worldwide genome sequencing projects has raised the following fundamental question to prominence in contemporary biological science: how do we compare two genomes [21, 38]?

Our aim here is not to define a similarity measure for each application field; we develop a general mathematical theory of similarity. To obtain evidence that the theory significantly addresses the question (and is not only theoretically satisfactory), we test it on real-world applications in a wide range of fields. Thus, we first present a new theoretical approach to a wide class of similarity metrics; show that the "normalized information distance" (possibly not in the class) is a metric, and prove that it is universal in the sense that this single metric uncovers all similarities simultaneously that the metrics in the class uncover a single similarity apiece. It is well-known that when a pure mathematical theory is applied to the real world, for example in hydrodynamics or in physics in general, we can in applications only approximate the theoretical ideal. But the theory gives a framework for the applied science. With this in mind, we demonstrate that the new universal similarity metric works well on concrete examples in very different application fields—the first completely automatic construction of the phylogeny tree based on whole mitochondrial genomes, and a completely automatic construction of a language tree for over 50 Euro-Asian languages. Other applications we have performed, not reported here, are detecting plagiarism in student programming assignments [34], and phylogeny of chain letters in [5].

Related Work: Preliminary applications of the current approach were tentatively reported to the biological community [22]—using an initial and partially improper metric. That work, and the present paper, is based on information distance [4], a single metric that captures in an appropriate sense every effective metric: effective versions of Hamming distance, Euclidean distance, edit distances, Lempel-Ziv distance, and the sophisticated distances introduced in [10, 27]. Subsequent work in the linguistics setting, [2, 3], used compression-based methods to infer a language tree from different-language text corpora, as well as to authorship attribution on basis of text corpora. Their methods approximate a certain type of empirical relative entropy—entropy à la Shannon is always zero since they deal with individual objects only. Even though the authors refer to the information distance in [25], the actual method is essentially ad-hoc without supporting theory, and based on "distances" violating metric requirements like symmetry and triangle inequality. The information distance studied in [24, 25, 4, 22], and subsequently investigated in [23, 17, 28, 30, 36], is "universal" and has other nice properties. This distance essentially says that the distance between two objects is the length
of the shortest program (or amount of energy) that is needed to transform the two objects into each other. But this distance is not proper to measure evolutionary sequence distance. For example, H. influenza and E. coli are two closely related sister species. The former has about 1,856,000 base pairs and the latter has about 4,772,000 base pairs. However, using the information distance of [4], one would easily classify H. influenza with a short (of comparable length) but irrelevant species simply because of length, instead of with E. coli. The information distance of [4] does not deal with relative distance. This is important because species may lose genes (by deletion) or gain genes (by duplication or insertion from external sources), relatively easily. Instead, deletion and insertion cost energy (proportional to the Kolmogorov complexity of deleted or inserted sequence) in the information distance of [4]. The paper [35] defined transformation distance between two species, and [16] defined compression distance. Both of these measures are essentially $K(x|y)$. Other than being asymmetric, they also suffer similar problems as the information distance of [4] as show in the above example. As far as the authors know, the idea of normalized metric is, surprisingly, not well studied. An exception is [39], which investigates normalized Euclidean metric and normalized symmetric-set-difference metric to account for relative distances rather than absolute ones, and it does so for much the same reasons as in the present work.

This Work: We develop a new general tool for phylogeny analysis and other applications: a general mathematical theory of similarity based on a special type of normalized metrics, and construct a universal normalized metric (the normalized information distance). This normalized information distance is the first universal effective similarity measure, and is an objective recursively invariant notion by the Church-Turing thesis. To evidence the significance of this theory, we apply it computationally in widely diverse real-world areas. (In the application, of course, we cannot compute the theoretical normalized information distance precisely, we have to do with approximations we can achieve.) In order to demonstrate universal applicability in an experimental setting, we use the diverse areas of (i) biochemical evolution studies, and (ii) natural language evolution. In area (i): In recent years, as the complete genomes of various species become available, it has become possible to do whole genome phylogeny (this overcomes the problem that different genes may give different trees). However, traditional phylogenetic methods on individual genes depended on multiple alignment of the related proteins and on the model of evolution of individual amino acids. Neither of these is practically applicable to the genome level. In absence of such models, a method which can compute the shared information between two sequences is useful because biological sequences encode information, and the occurrence of evolutionary events (such as insertions, deletions, point mutations, rearrangements, and inversions) separating two sequences sharing a common ancestor will result in the loss of their shared information. Our theoretical approach is used experimentally to create a fully automated and reasonably accurate software tool based on such a distance to compare two genomes. We demonstrate that a whole mitochondrial genome phylogeny of the Eutherians which confirms [8], can be reconstructed automatically from unaligned complete mitochondrial genomes by use of our software implementing (an approximation of) our theory. These experimental confirmations of the efficacy of our comprehensive approach contrasts with recent more specialized approaches such as [37] that have (and perhaps can) only be tested on small numbers of genes. They have not been experimentally tried on whole mitochondrial genomes that are, apparently, already numerically out of computational range. In area (ii) we fully automatically construct the language tree of 52 primarily Indo-European languages from translations of the "Universal Declaration of Human Rights"—leading to a grouping of language families largely consistent with current linguistic viewpoints. The main technical concepts in this work, distance metric, Kolmogorov complexity [25], information distance as in [4], are summarized below.

2 Preliminaries

Metric: Without loss of generality, a distance only needs to operate on sequences of 0's and 1's since any sequence can be represented by a binary string. Formally, a distance function $D$ with nonnegative real values, defined on the Cartesian product $X \times X$ of a set $X$ is called a metric on $X$ if for every $x, y, z \in X$:

- $D(x, y) = 0$ iff $x = y$ (the identity axiom);
- $D(x, y) + D(y, z) \geq D(x, z)$ (the triangle inequality);
- $D(x, y) = D(y, x)$ (the symmetry axiom).

A set $X$ provided with a metric is called a metric space. For example, every set $X$ has the trivial discrete metric $D(x, y) = 0$ if $x = y$ and $D(x, y) = 1$ otherwise. All distances in this paper are defined on the set $X = \{0, 1\}^*$ and satisfy the metric conditions sometimes up to an additive vanishing constant term. In our search for the proper definition of the distance between two, not necessarily equal length, binary strings, a natural choice is the length of the shortest program that can transform either string into the other one—both ways. This distance is known as information distance, which is one of the main concepts in this work and which we will discuss in detail below. However, such a distance measures an absolute distance, and we are more interested in a relative one. For example, if two strings of length $10^6$ have distance 1000, then we are inclined to think that those strings are relatively more similar than two strings of length 1000 that have that distance and consequently are 100% different. Therefore, we want to normalize the information distance into a normalized distance. We do this by dividing it by the greater of the two lengths of the shortest programs that compute the strings concerned from scratch.

Kolmogorov Complexity: An introduction, details, and proofs of the theory of Kolmogorov complexity can be found in the text [25]. Here we recall some basic notation and facts. We write $\text{string}$ to mean a finite binary string. Other finite objects can be encoded into strings in natural ways. The set of strings is denoted by $\{0, 1\}^*$.
The Kolmogorov complexity, or algorithmic entropy, $K(x)$ of a string $x$ is the length of a shortest binary program to compute $x$ on an appropriate universal computer—such as a universal Turing machine. (It is equivalent to consider the length of the shortest binary program to compute $x$ in a universal programming language such as LISP or Java.) The functions $K(\cdot)$ and $K(\cdot|\cdot)$, though defined in terms of a particular machine model, are machine-independent up to an additive constant and acquire an asymptotically universal and absolute character through Church’s thesis, from the ability of universal machines to simulate one another and execute any effective process. Intuitively, $K(x)$ represents the minimal amount of information required to generate $x$ by an algorithm, [20]. $x^*$ denotes a shortest program for $x$ (if there is more than one of them then the first one in standard enumeration), and hence $|x^*| = K(x)$. The conditional Kolmogorov complexity $K(x|y)$ of $x$ relative to $y$ is defined similarly as the length of a shortest program to compute $x$ if $y$ is furnished as an auxiliary input to the computation. We use the notation $K(x, y)$ for the length of a shortest binary program that prints out $x$ and $y$ and a description how to tell them apart.

**DEFINITION 2.1.** A real-valued function $f(x, y)$ is upper semi-computable if there exists a rational-valued recursive function $g(x, y, t)$ such that (i) $g(x, y, t + 1) \leq g(x, y, t)$, and (ii) $\lim_{t \to \infty} g(x, y, t) = f(x, y)$. It is lower semi-computable if $-f(x, y)$ is upper semi-computable, and it is computable if it is both upper- and lower semi-computable.

It is easy to see that the functions $K(x)$ and $K(y|x^*)$ (and under the appropriate interpretation also $x^*$) are upper semi-computable, and it is easy to prove that they are not computable. The conditional information contained in $x^*$ is equivalent to that in $(x, K(x))$. The information in $y$ about $x$ is defined as $I(x : y) = K(x) - K(x|y^*)$. A deep, and very useful, result [13] shows that, up to additive constant terms $I(x : y) = I(y : x)$, that is

$$K(x) + K(y|x^*) = K(y) + K(x|y^*).$$

**Information Distance:** The information distance, [4], is the length of a shortest binary program that computes $x$ from $y$ as well as computing $y$ from $x$. Being shortest, such a program should take advantage of any redundancy between the information required to go from $x$ to $y$ and the information required to go from $y$ to $x$. The program functions in a catalytic capacity in the sense that it is required to transform the input into the output, but itself remains present and unchanged throughout the computation. A principal result of [4] shows that, up to an additive logarithmic term, the information distance equals

$$E(x, y) = \max\{K(y|x), K(x|y^*)\}$$

Because of the upper semi-computability of the conditional complexities, the information distance is also upper semi-computable. (It is very important here that the time of computation is completely ignored: this is why this result does not contradict the idea of one-way functions.)

**3 Normalized Distance**

In defining a class of acceptable metrics we want to exclude unrealistic distance metrics like $f(x, y) = \frac{1}{2}$ for every pair $x \neq y$, by restricting the number of objects within a given distance of an object. As in [4] we only consider upper semi-computable distances $D(x, y)$ satisfying the density condition

$$\sum_{y \neq x} 2^{-D(x, y)} \leq 1.$$  

As remarked above, large objects (in the sense of long strings) that differ by a tiny part are intuitively closer than tiny objects that differ by the same amount. Therefore, we normalize the distance metric: Let $D(x, y)$ be an upper semi-computable distance satisfying the density condition (3.3). Let $n$ be a function such that $d_n, D(x, y) = n(D, x, y)$ has values in $[0, 1]$ and satisfies the normalization condition

$$\sum_{y \neq x} 2^{-d_n, D(x, y)} K(x) \leq 1.$$  

Then, with $K(x) = k$ and $d \in [0, 1]$, we have

$$\sum_{y \neq x} 2^{-d_n, D(x, y)} K(x) \leq 2^{-dk}.$$  

For suppose the contrary: Starting from (3.4) we obtain a contradiction:

$$1 \geq \sum_{y \neq x} 2^{-d_n, D(x, y)} K(x) \geq 2^{-dk} > 2^{dk}2^{-dk} = 1.$$  

**DEFINITION 3.1.** A normalized distance or similarity distance is a metric $m(x, y)$ that takes values in $[0, 1]$ (up to an additive term that vanishes with growing $\max\{K(x), K(y)\}$) for all $x, y$, and satisfies the density constraint (3.5).

**4 Normalized Information Distance**

Clearly, unnormalized information distance (2.2) is not a proper evolutionary distance measure. Consider three species: *E. coli*, *H. influenza*, and some arbitrary bacteria *X* of similar length as *H. influenza*, but not related. Information distance $d$ would have $d(X, H.influenza) < d(E.coli, H.influenza)$, simply because of the length factor. It would put two long and complex sequences that differ only by a tiny fraction of the total information as dissimilar as two short sequences that differ by the same absolute amount and are completely random with respect to one another.

In [22] we considered as first attempt at a normalized information distance:

**DEFINITION 4.1.** Given two sequences $x$ and $y$, define the function $d_s(x, y)$ by

$$d_s(x, y) = \frac{K(x | y^*) + K(y | x^*)}{K(x, y)}.$$
Writing it differently, using (2.1),
\[ d_s(x, y) = 1 - \frac{K(x) - K(x \mid y^*)}{K(x, y)}, \]
where \( K(x) - K(x \mid y^*) \) is the mutual information \( I(y : x) \).
This distance satisfies the triangle inequality, up to a small error term, and universality (below), but only within a factor 2. Mathematically more precise and satisfying is the distance:

**DEFINITION 4.2.** Given two sequences \( x \) and \( y \), define the function \( d(x, y) \) by
\[
(4.7) \quad d(x, y) = \max\{K(x \mid y^*), K(y \mid x^*)\}.
\]

**REMARK 4.3.** Several natural alternatives for the denominator turn out to be wrong:
(a) Divide by the length. Then, firstly we do not know which of the two length involved to divide by, possibly the sum or maximum, but furthermore the triangle inequality and the universality (domination) properties are not satisfied.
(b) In the \( d \) definition divide by \( K(x, y) \). Then one has \( d(x, y) = \frac{1}{2} \) whenever \( x \) and \( y \) are random (have maximal Kolmogorov complexity) relative to one another. This is improper.
(c) In the \( d_s \) definition dividing by length does not satisfy the triangle inequality.

There is a natural interpretation to \( d(x, y) \): If \( K(y) \geq K(x) \) then we can rewrite
\[
d(x, y) = \frac{K(y) - I(x : y)}{K(y)},
\]
where \( I(x : y) \) is the information in \( y \) about \( x \) satisfying the symmetry property \( I(x : y) = I(y : x) \) up to a logarithmic additive error by (2.1). That is, the ratio \( d(x, y) \) between \( x \) and \( y \) is the number of bits of information that is not shared between the two strings per bit of information that could be maximally shared between the two strings.

It is clear that \( d(x, y) \) is symmetrical and satisfies the identity axiom:
\[
d(x, x) = O(1/K(x)).
\]
To show that it is a distance metric it remains to prove the triangle inequality.

**LEMMA 4.4.** \( d(x, y) \) almost satisfies the weak triangle inequality, that is, \( d(x, y) \leq d(x, z) + d(z, y) \) up to an additive error term of \( O(1/\max\{K(x), K(y), K(z)\}) \).

*Proof. Case 1:* Suppose \( K(z) \leq \max\{K(x), K(y)\} \). In [14], the following “directed triangle inequality” was proved: For all \( x, y, z \), up to an additive constant term,
\[
(4.8) \quad K(x \mid y^*) \leq K(z \mid y^*) \leq K(x \mid z^*) + K(z \mid y^*).
\]
Dividing both sides of the triangle inequality by \( \max\{K(x), K(y)\} \),
\[
\frac{\max\{K(x \mid y^*), K(y \mid x^*)\}}{\max\{K(x), K(y)\}} \leq \frac{\max\{K(x \mid z^*), K(z \mid x^*)\} + \max\{K(x \mid y^*), K(y \mid z^*)\}}{\max\{K(x), K(y)\}},
\]
up to an additive term \( O(1/\max\{K(x), K(y), K(z)\}) \). Replacing \( K(x) \) or \( K(y) \) in the denominator of the first term of the right-hand side by \( K(z) \) only increase the right-hand side (again, because of the assumption).

**Case 2:** Suppose \( K(z) = \max\{K(x), K(y), K(z)\} \). Further assume that \( K(x) \geq K(y) \) (the remaining case is symmetrical). Then, using the symmetry of information to determine the maxima, we also find \( K(z \mid x^*) \geq K(z \mid z^*) \) and \( K(z \mid y^*) \geq K(y \mid z^*) \). Then the maxima in the terms of the equation \( d(x, y) \leq d(x, z) + d(y, z) \) are determined, and our proof obligation reduces to:
\[
(4.9) \quad \frac{K(x \mid y^*)}{K(x)} \leq \frac{K(z \mid x^*) + K(z \mid y^*)}{K(z)},
\]
up to an additive term \( O(1/K(z)) \). To prove (4.9) we proceed as follows:

Applying the triangle inequality (4.8) and dividing both sides by \( K(x) \), we have
\[
\frac{K(x \mid y^*)}{K(x)} \leq \frac{K(z \mid x^*) + K(z \mid y^*) + O(1)}{K(z)}.
\]
The left-hand side is \( \leq 1 \).

**Case 2.1:** The right-hand side is \( \leq 1 \). Setting \( K(z) = K(x) + \Delta, \) and first adding \( \Delta = K(z) - K(x) = K(z \mid x^*) - K(x \mid z^*) + O(1) \) to both the nominator and the denominator of the first term in the right-hand side, and subsequently using (2.1) to obtain \( K(x \mid z^*) + \Delta = K(z \mid z^*) + O(1) \),
\[
\frac{K(x \mid y^*)}{K(x)} \leq \frac{K(x \mid z^*) + K(z \mid y^*) + \Delta + O(1)}{K(x)},
\]
which was what we had to prove.

**Case 2.2:** The right-hand side is \( \geq 1 \). We proceed like in Case 1, and add \( \Delta \) to both nominator and denominator. Although now the right-hand side decreases, it must still be \( \geq 1 \). This proves Case 2.2.

Clearly, \( d(x, y) \) takes values in the range \([0, 1 + O(1/\max\{K(x), K(y)\})]\). To show that it is a normalized distance, it is left to prove the normalization condition:

**LEMMA 4.5.** The function \( d(x, y) \) satisfies the normalization condition (3.5).

*Proof.* Assume that \( K(y) \geq K(x) \) (the other case is symmetrical). Then, by (2.1) we also have \( K(y \mid x^*) \geq K(x \mid y^*) \) up to an additive constant term, and rewriting \( d(x, y) = K(y \mid x^*)/K(y) \leq d \) and \( K(y) \leq k \), we obtain up to an additive constant term \( K(y \mid x^*) \leq dK(y) \leq dk \). That is, there are at
most $2^{dk}$ binary programs to obtain a $y$ from $x^*$ (recall that we assume $K(y) \geq K(x)$), and hence at most that many such $y$.

Since we have shown that $d(x, y)$ is a distance metric, takes values in $[0,1]$, (up to vanishing additive error terms) and satisfies the normalization condition, it follows:

**Theorem 4.1.** The function $d(x, y)$ is a normalized information distance.

### 5 Universality

We now show that $d(x, y)$ is universal in the sense that it incorporates every remotely computable type of similarity in the following sense: If two objects are similar in normalized information in some computable sense, then they are at least that similar in the $d(x, y)$ sense. We prove this by demonstrating that $d(x, y)$ is smaller than every other normalized distance in a wide class—so wide that it will capture everything that can be remotely of interest. The function $d(x, y)$ itself, being a ratio between two maxima of pairs of upper semi-computable functions, is not itself upper semi-computable. (It is easy to see that this is likely, but a formal proof is difficult.) In fact, $d(x, y)$ has ostensibly only a weaker computability property: Call a function $f(x, y)$ computable in the limit if there exists a rational-valued recursive function $g(x, y, t)$ such that $\lim_{t \to \infty} g(x, y, t) = f(x, y)$. Then $d(x, y)$ is in this class. It can be shown (in the full version) that this is precisely the set of functions that are Turing-reducible to the halting set. While $d(x, y)$ is not upper semi-computable, it captures all similarities represented by the normalized metrics in the class concerned, which should suffice as a theoretical basis for all practical purposes.

**Theorem 5.1.** The normalized information distance $d(x, y)$ dominates every upper semi-computable normalized distance $f(x, y)$ up to a vanishing additive term: $d(x, y) \leq f(x, y) + O(\log k)/k$, where $k = \max\{K(x), K(y)\}$.

**Proof.** Fix a normalized distance $f(x, y)$ and assume $f(x, y) = d$. By the normalization condition we have that, given $x$, the number of $y$, such that $f(x, y) \leq d$ and $\max\{K(x), K(y)\} = k$, is upper bounded by $2^{dk}$. Hence, for fixed $x^*$ and $k$ we can recursively enumerate the $y$ for which $f(x, y) \leq d$ and $K(y) \leq k$, and every $y$ can be described by its index of length $dk$ in this enumeration. Since the Kolmogorov complexity is the length of the shortest effective description, the binary length of the index must at least be as large as the Kolmogorov complexity, which yields $K(y \mid x^*, k) = K(y \mid x, K(x), k) \leq dk$. That is, since we can provide both $k$ and $K(x)$ in $O(\log k)$ bits, $K(y \mid x^*) \leq dk + O(\log k)$.

Substitution gives:

$$d(x, y) = \frac{K(y \mid x^*)}{K(y)} \leq \frac{dk + O(\log k)}{k} = f(x, y) + O\left(\frac{\log k}{k}\right).$$

The other case, $K(x) > K(y)$ (so $K(x) = k$) gives:

$$d(x, y) = \frac{K(x \mid y^*)}{K(x)} \leq \frac{dk + O(\log k)}{k} = f(x, y) + O\left(\frac{\log k}{k}\right).$$

### 6 Application to Whole Mitochondrial Genome Phylogeny

Nothing is more ideal than DNA sequences to test our theory. We will use whole mitochondrial DNA genomes of 20 mammals and the problem of Eutherian orders to make a comprehensive examination of our measures. The problem we consider is this: It has been debated in biology which two of the three main groups of placental mammals are more closely related: Primates, Ferungulates, and Rodents. This is because the maximum likelihood method gives (Ferungulates, (Primates, Rodents)) grouping for half of the proteins in mitochondrial genome, and (Rodents, (Ferungulates, Primates)) for the other half [8]. In [8], Cao et al. aligned 12 concatenated mitochondrial proteins taken from the following species: rat (Rattus norvegicus), house mouse (Mus musculus), grey seal (Halichoerus grypus), harbor seal (Phoca vitulina), cat (Felis catus), white rhino (Ceratotherium simum), horse (Equus caballus), finback whale (Balaenoptera physalus), blue whale (Balaenoptera musculus), cow (Bos taurus), gibbon (Hylobates lar), gorilla (Gorilla gorilla), human (Homo sapiens), chimpanzee (Pan troglodytes), pygmy chimpanzee (Pan paniscus), orangutan (Pongo pygmaeus), Sumatran orangutan (Pongo pygmaeus abelii), using opossum (Didelphis virginiana), wallaroo (Macropus robustus) and platypus (Ornithorhynchus anatinus) as the outgroup, and built the maximum likelihood tree. The resulting phylogeny supports the currently accepted grouping (Rodents, (Primates, Ferungulates)).

#### 6.1 Alternative Approaches: Before applying our theory to further confirm this hypothesis, we first examine the alternative approaches, in addition to that of [8]. The mitochondrial genomes of the above 20 species were obtained from GenBank. Each is about 18k bases.

**K-mer Statistic:** In the early years, researchers experimented on using G+C contents or slightly more general k-mers (or Shannon block entropy) to classify DNA sequences (in particular S. Wildman at Stanford). This approach uses the statistics of length $k$ substrings in a genome and the phylogeny is constructed accordingly. To re-examine this approach, we performed simple experiments: Consider all length $k$ blocks in each mtDNA, for $k = 1, 2, \ldots, 10$. There are $l = (4^{11} - 1)/3$ different sequences (some may not appear in an mtDNA). We computed their number of occurrences in each mtDNA, obtaining a vector of length $l$ for each mtDNA.

For two such vectors (representing two mtDNAs) $p, q$, their distance is computed as $d(p, q) = \sqrt{(p - q)^T(p - q)}$. Using neighbor joining [32], the resulting tree is the one given in Figure 1. Using the hypercleaning method [7], we obtain equally absurd results. Similar experiments were repeated for size $k$ blocks alone (for $k = 10, 9, 8, 7, 6$), without much improvement.
Gene Order: In [6] the authors propose to use the order of genes to infer the evolutionary history. This approach does not work for closely related species such as our example where all genes are in the same order in the order of genes to infer the evolutionary history. This mammalian mtDNA sequences using frequency of k-

Figure 1: The evolutionary tree built from complete mammalian mtDNA sequences using frequency of k-

Gene Content: Gene content method, proposed in [12, 33], compares number of genes two species share divided by total number of genes. While this approach does not work here due to the fact that all 20 mammalian mitochondrial genomes share exactly the same genes, notice the similarity of gene content formula and our general formula.

Rearrangement Distance: Reversal and rearrangement distances in [19, 18, 29] compare genomes using other partial genome information such as number of reversals or translocations. These operations also do not appear in our mammalian mitochondrial genomes, hence the method again is not proper for our application.

Transformation Distance or Compression Distance: The transformation distance proposed in [35] and compression distance proposed in [16] are essentially defined as $K(x|y)$ which is asymmetric, and so, is not a distance. The measure $K(x|y)$ produces a wrong tree with one of the marsupials mixed up with ferungulates (the tree is not shown here).

6.2 The Present Approach We have shown that $d$ (and up to a factor 2 also $d_s$) is universal among a wide class of computable normalized information measures. However the generality of $d$ and $d_s$ comes at the price of noncomputability: Kolmogorov complexity is not computable but upper semi-computable, Section 2, and $d$ and $d_s$ are (likely to be) not even that. Nonetheless, we can try to approximate the spirit of $d$ and $d_s$ at various levels of precision. Now it is clear how to upper semi-compute the unconditional complexities involved. With respect to the conditional complexities, by (2.1) we have $K(x | y) = K(x, y) - K(y)$ (up to an additive constant), and it is easy to see that $K(x, y) = K(xy)$ up to additive logarithmic precision. (To retrieve $(x, y)$ we need to encode the separator between the binary programs for $x$ and $y$.)

k-mers According to the New Measures: We have shown that using k-mer statistics alone does not work well. However, let us now combine the k-mer approach with the new measures. Consider the length-k substrings of the DNA sequence as words of the sequence. We denote the number of distinct, possibly overlapping, k-length words in a sequence $x$ by $N(x)$. With $k$ large enough, at least $\log_a(n)$, where $a$ is the cardinality of the alphabet and $n$ the length of $x$, we use $N(x)$ as a rough approximation to $K(x)$. We justify this by the pragmatic observation that, because the genomes evolve by duplications, rearrangements and mutations, [31], it can be argued that it is appropriate to use $N(x)$ to very roughly estimate for $K(x)$ in case $x$ is a genome. Define $N(x|y)$ as $N(xy) - N(y)$. Given two sequences $x$ and $y$, following the definition of $d$, (4.7), the distance between $x$ and $y$ can be defined as

$$d'(x, y) = \frac{\max\{N(x|y), N(y|x)\}}{\max\{N(x), N(y)\}}.$$  

Similarly, following $d_s$, (4.6) we can also define another distance using $N(x)$,

$$d^*(x, y) = \frac{N(x|y) + N(y|x)}{N(xy)}.$$  

Using $d'$ and $d^*$, we computed the distance matrices for the 20 mammal mitochondrial DNAs. Then we used hyper-Cleaning [7] to construct the phylogenies for the 20 mammals. Using either of $d'$ and $d^*$, we were able to construct the tree correctly when $8 \leq k \leq 13$, as in Figure 3. A tree constructed with $d'$ for $k = 7$ is given in Figure 2. We note that the opossum and a few other species are misplaced. The tree constructed with $d^*$ for $k = 7$ is very similar, but it correctly positioned the opossum.

Spaced k-mers According to the New Measures In methods for doing DNA homology search, a pair of identical words, each from a DNA sequence, is called a “hit”. Hits have been used as “seeds” to generate a longer match between the two sequences. We note that $N(x|y)$ is the number of distinct words that are in $x$ and not in $y$, the more hits the two sequences have, the smaller the $N(x|y)$ and $N(y|x)$ are. Therefore, the previous two distances can also be interpreted as a function of the number of hits, each of which indicates some mutual information of the two sequences.

As noticed by the authors of [26], though it is difficult to get the first hit (of consecutive k letters) in a region, it only requires one more base match to get a second hit overlapping the existing one. This makes it inaccurate to attribute the same amount of information to each of the hits. For this reason, we also tried to use the “spaced model” introduced in [26] to compute our distances. A length-L, weight-k spaced model is a 0-1 string of length $L$ and having $k$ 1s. We overlap such a model with the DNA sequence at
Each of the positions in the DNA sequence, and take out the $k$ bases covered by the 1s to form a length-$k$ word. The number of those distinct words is then used to define the distances $d'$ and $d^*$ in Formula (6.2) and (6.2).

We applied the new defined distances to the 20 mammal data. The performance is slightly better than the performance of the distances defined in (6.2) and (6.2). The modified $d'$ and $d^*$ can correctly construct the mammal tree when $7 \leq k \leq 13$ and $6 \leq k \leq 13$, respectively.

Compression: To achieve the best approximation of Kolmogorov complexity, and hence most confidence in the approximation of $d_*$ and $d$, we use a new version of the GenCompress program, [9], which achieves the currently best compression ratios for benchmark DNA sequences. GenCompress finds approximate matches (hence edit distance becomes a special case), approximate reverse complements, among other things, with arithmetic encoding when necessary. Online service of GenCompress is at UCSB Bioinformatics Lab website: http://cytosine.cs.ucsb.edu:8080/.

We computed $d(x, y)$ between each pair of mtDNA $x$ and $y$, using GenCompress to heuristically approximate $K(x|y)$, $K(x)$, and $K(x, y)$, and constructed a tree (Figure 3) using the neighbor joining [32] program in the MOLPHY package [1]. The tree is identical to the maximum likelihood tree of Cao, et al. [8]. For comparison, we used the hypercleaning program [7] and obtained the same result. The phylogeny in Figure 3 re-confirms the hypothesis of (Rodents, (Primates, Feringulates)).

To further assure our results, we have extracted only the coding regions from the mtDNAs of the above species, and performed the same computation. This resulted in the same tree.

Evaluation: This new method for whole genome comparison and phylogeny does not require gene identification nor any human intervention, in fact, it is totally automatic.

It is mathematically well-founded being based on general information theoretic concepts. It works when there are no agreed upon evolutionary models, as further demonstrated by the successful construction of a chain letter phylogeny [5] and when individual gene trees do not agree (Cao et al., [8]) as is the case for genomes. Next step would be to apply this method to larger genomes such as cpDNA and bacteria genomes.

7 The Language Tree

Normalized information distance is a totally general universal tool, not restricted to a particular application area. We show that it can also be used to successfully classify natural languages. Let us borrow from biology the “nature” (acquired by genetic mixture) versus “nurture” (acquired in the life of the individual) terminology. Any language tree built by only analyzing contemporary natural text corpora is partially corrupted by historical “nurture” contaminations. While according to Darwinism the genomes only change by inheritance (nature), languages acquire their characteristic by descent but also by interaction (nurture). Thus, while English is a Germanic Anglo-Saxon language, it has absorbed a great deal of French-Latin components. Similarly, Hungarian, often considered a Finno-Ugric language, which consensus currently happens to be open to debate in the linguistic community, is known to have absorbed many Turkish and Slavic components. Thus, an automatic construction of a language tree based on contemporary text corpora, exhibits current linguistic relations (based on both nature and nurture) which do not necessarily coincide completely with the historic language family tree (based on nature). According to a linguistic expert, only vocabulary is normally borrowed between languages, and inflectional morphology is the best indicator of linguistic descent. This may be the most important factor distorting the results. The misclassification of English as Romance language must have something to do
Figure 4: The language classification tree, built from "The Universal Declaration of Human Rights" [15] using approximated normalized information distance, $d_n$-version (6.2), and neighbor joining. This is a rooted tree, using Basque [Spain] as outgroup. The branch lengths are not proportional to the actual distances in the distance matrix.

Figure 5: The language classification tree, built from "The Universal Declaration of Human Rights" [15] using approximated normalized information distance, $d_n$-version (4.7), and the Fitch-Margoliash method. This is a rooted tree using Basque [Spain] as outgroup. The branch lengths are not proportional to the actual distances in the distance matrix.

with the fact that the English vocabulary in the Universal Declaration of Human Rights, being nonbasic in large part, is Latinate in large part. It also accounts for the misclassification of Maltese, an Arabic dialect with lots of Italian loan words, as Romance. Having voiced these caveats, the result of our automatic experiment in language tree reconstruction is rather encouraging.

Our method improves the results of [2], using a linguistic corpus of "The Universal Declaration of Human Rights" [15] in 52 languages. The previous effort [2] used an asymmetric measure based on relative entropy, and the full matrix of the pair-wise measures between all 52 languages, to build a language classification tree. This resulted in some inconsistencies, such as English being isolated between Romance and Celtic languages, Romani-balkan and Albanian being isolated, and Hungarian (possibly a Finno-Ugrian language) being grouped with Turkish and Uzbek. The (rooted) trees resulting from our experiments (using Basque as outgroup) seem more correct. We use Basque as outgroup since linguists regard it as a language unconnected to other languages. In order to test the potential of normalized information distance $d$ in classifying natural languages, a similar experiment was performed. First, transform each UNICODE character in the language text into an ASCII character by removing its vowel flag if necessary. Secondly, a LZ-type algorithm gzip is appropriate to compress these language sequences of sizes not exceeding the length of sliding window gzip use (32 kilobytes), and compression results can be used to approximate their Kolmogorov complexity. Instead of a complicated and less rigorous method to approximate relative entropy used in [2], we simply zip-compress several sequences that involve calculating the normalized information metric (both (6.2) and (4.7)) based on each pair of language
sequences. In the last step, we applied the $d_n$-metric (6.2) with the neighbor-joining package to obtain Figure 4. Even better worked applying the $d$-metric (4.7) with the Fitch-Margoliash method [11] in the package PHYLIP [1]); the resulting language classification tree is given in Figure 5. We note that all the main linguistic groups can be successfully recognized, which includes Romance, Celtic, Germanic, Ugro-Finnic, Slavic, Baltic, Altaic as labeled in the figure.

8 Conclusion

We developed a mathematical theory of similarity distances and shown that there is a universal similarity distance: the normalized information distance. This distance uncovers all computable similarities, and therefore estimates an evolutionary or relation-wise distance on strings. It has been shown to be applicable to whole genomes, but as well to chain letters (5), not included here), to test students source code for plagiarism (34), not included here), and to build a large language family tree from text corpora. It is perhaps useful to point out that the results reported in the figures were obtained at the very first runs and have not been selected by appropriateness from several trials. From the theory point-of-view we have obtained a general mathematical theory forming a solid framework spawning practical tools applicable in many fields. Based on the noncomputable notion of Kolmogorov complexity, the normalized information distance can only be approximated in an ad hoc manner, that is, without speed of convergence guarantees. Even so, the fundamental rightness of the approach is evidenced by the remarkable success (agreement with known phylogeny in biology) of the evolutionary trees obtained and the building of language trees. From the applied side of genomics our work gives the first fully automatic generation of whole genome mitochondrial phylogeny; in computational linguistics it presents a fully automatic way to build language trees and determine language families.

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