Power of the power-laws: lessons from unification of small and large time scales for evolution

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We develop a “unified” model that describes both “micro” and “macro” evolutions within a single theoretical framework. The eco-system is described as a dynamic network; the population dynamics at each node of this network describes the “micro”-evolution over ecological time scales (i.e., birth, ageing and natural death of individual organisms) while the appearance of new nodes, the slow changes of the links and the disappearance of existing nodes accounts for the “macro” evolution over geological time scales (i.e., the origination, evolution and extinction of species). In contrast to several earlier claims in the literature, we observe strong deviations from power law in the regime of long life times where the statistics is, usually, poor.

The recent surge in the modeling of biological evolution and extinction of species, using the concepts and techniques of statistical physics, has been stimulated partly by the claims (see \cite{1,2,3,4} for reviews) that the statistical distributions of several quantities associated with the extinction of species follow power laws. However, almost all of these models focus only on the “macro” evolution (i.e., the evolution of species on geological time scales). Neither the birth, ageing and, eventually, the death of the individual organisms nor the detailed population dynamics make explicit appearance in these theoretical descriptions. On the other hand, in reality, a species becomes extinct when its entire population is wiped out.

Therefore, we develop a “unified” model of an eco-system that describes both “micro” and “macro” evolutions. The eco-system is described as a dynamic network. The “micro”-evolution over ecological time scales, i.e., birth, growth (ageing) and natural death of individual organisms is described by the dynamics within each node \cite{5}. The network itself evolves slowly with time; over sufficiently long time scales populations of some species would drop to zero, indicating their extinction, and the corresponding nodes would be deleted from the network. On the other hand, appearance of new nodes, together with their own population of individual organisms, signals origination of new species. In addition, the links of the network also change slowly to capture the adaptive evolution of the species by random mutations over geological time scales.

\textbf{The dynamic network:} At any arbitrary instant of time $t$ the model consists of $N(t)$ species each of which may be represented by one of the $N$ nodes of a dynamic network; the total number of nodes is not allowed to exceed $N_{\text{max}}$. Our model allows $N(t)$ to fluctuate with time over the range $1 \leq N(t) \leq N_{\text{max}}$. The population (i.e., the total number of organisms) of a given species, say, $i$ at any arbitrary instant of time $t$ is given by $n_i(t)$. The limited availability of resources, other than food, in the eco-system imposes an upper limit $n_{\text{max}}$ of the allowed population of each species. Thus, the total number of organisms $n(t)$ at time $t$ is given by $n(t) = \sum_{i=1}^{N(t)} n_i(t)$. Both $N_{\text{max}}$ and $n_{\text{max}}$ are time-independent in the model.

\textbf{The interactions:} Prey-predator interactions are captured through the matrix $J$. The influence of species $j$ on species $i$ is given by $J_{ij}$; in general, $J_{ij} \neq J_{ji}$. The only restriction we impose initially on the elements of $J$ is that $J_{ii} = 0$, i.e., none of the organisms preys on any other member of the same species. Since in all practical situations, food webs specify only the sign of $J_{ij}$ we allow the off-diagonal elements of $J_{ij}$ to take only the values +1 and −1. Thus, $J_{ij} = 1 = -J_{ji}$ indicates that $j$ is the prey and $i$ is the predator. Similarly, the situations $J_{ij} = -1 = J_{ji}$ and $J_{ij} = 1 = J_{ji}$ correspond, respectively, to \textit{competition} and \textit{cooperation} between the species $i$ and $j$. We assign the values +1 or −1 to the off-diagonal elements of $J$ randomly with equal probability in the initial state of the eco-system \cite{6}. However, our model can be easily generalized to take into account any other architecture of food webs \cite{7,8}.

We now argue that the matrix $J$ accounts not only for the \textit{inter}-species interactions but also \textit{intra}-species interactions. First of all, note that if $J_{ij} > 0$, then the species $j$ is a prey of the species $i$ if, simultaneously, $J_{ji} < 0$ whereas the species $j$ cooperates with $i$ if, simultaneously, $J_{ij} > 0$ and $J_{ji} > 0$. Therefore, if $J_{ij} > 0$, the quantity $(J_{ij} - J_{ji})/2$ is unity if the species $j$ is a prey of the species $i$, but it vanishes if the species $i$ and $j$ mutually cooperate. Similarly, if $J_{ij} < 0$, the quantity $-(J_{ij} - J_{ji})/2$ is unity if the species $j$ is a predator of $i$, but it vanishes if the species $i$ and $j$ compete against each other. Now, consider the two sums

\begin{equation}
S_i^\pm = \pm \sum_{j=1}^{N} \frac{(J_{ij}^\pm - J_{ji})}{2} n_j
\end{equation}

where the superscript $\pm$ on $J_{ij}$ indicates that the sum is restricted to only the positive (negative) elements $J_{ij}$. The sum $S_i^+$ is a measure of the total food currently available to the $i$-th species whereas $-S_i^-$ is a measure of the total population of the $i$-th species that would be, at the same time, consumed as food by its predators. If the food available is less than the requirement, then some organisms of the species $i$ will die of starvation, even if none of them is killed by any predator. This way the
matrix $J$ can account for the shortfall in the food supply and the consequent competition among the organisms of the species $i$.

The collective characteristics of species: The age of an arbitrary individual organism, say, $\alpha$ of the species $i$ at time $t$ is denoted by the symbol $X(i, \alpha; t)$. In our model each species $i$ is collectively characterized by the following:

(i) the minimum reproduction age $X_{\text{rep}}(i)$, (ii) the birth rate $M(i)$, (iii) the maximum possible age $X_{\text{max}}(i)$, and (iv) the elements $J_{ij}$ and $J_{ji}$ ($j = 1, 2, ..., N$). An individual of the $i$-th species can reproduce only after attaining the age $X_{\text{rep}}(i)$. Whenever an organism of $i$-th species gives birth to offsprings, $M(i)$ of these are born simultaneously. None of the individuals of the $i$-th species can live longer than $X_{\text{max}}(i)$. Thus, even if an individual manages to escape its predators, it cannot live longer than $X_{\text{max}}(i)$ because of “natural death” caused by ageing.

The dynamics of the eco-system: The state of the system is updated in discrete time steps as follows:

Step I - Birth: Assuming, for the sake of simplicity, the reproductions to be asexual, each individual organism $\alpha$ ($\alpha = 1, ..., n_i(t)$) of the species $i$ ($i = 1, 2, ..., N(t)$) is allowed to give birth to $M(i; t)$ offsprings at every time step $t$ with probability (per unit time) $p_b(i, \alpha; t)$ which is non-zero only when the individual organism age $X(i, \alpha; t) > X_{\text{rep}}(i; t)$.

Step II - Natural death: At any arbitrary time step $t$ the probability (per unit time) of “natural” death (due to ageing) of an individual organism $\alpha$ of species $i$ is $p_d(i, \alpha; t)$.

Step III - Mutation: With probability $p_{\text{mut}}$ per unit time, all the species simultaneously re-adjust one of the incoming interactions $J_{ij}$ by assigning it a new value of either +1 or −1 with equal probability.

Step IV - Starvation death and killing by prey: If $n_i - S^+$ is larger than $S^-$ then food shortage will be the dominant cause of premature death of a fraction of the existing population of the species $i$. On the other hand, if $S^- > n_i - S^+$, then a fraction of the existing population will be wiped out primarily by the predators. In order to capture these phenomena, at every time step $t$, in addition to the natural death due to ageing, a further reduction of the population by

$$C \max(S^-, n_i - S^+)$$

is implemented where $n_i(t)$ is the population of the species $i$ that survives after the natural death step above. $C$ is a constant of proportionality that basically sets the time scale of the population dynamics. If implementation of these steps makes $n_i(t) \leq 0$, species $i$ becomes extinct.

Step V - Speciation: At each time step, the niches (nodes) left empty by extinction of species are re-filled by new species, with probability $p_{\text{sp}}$. All the simultaneously re-filled nodes of the network originate from one common ancestor which is picked up randomly from among the surviving species. All the interactions $J_{ij}$ and $J_{ji}$ of the new species are identical to those of their common ancestor; each new species, however, either competes or cooperates with its ancestor species. The characteristic parameters $X_{\text{max}}$, $X_{\text{rep}}$, $M_j$ of each of the new species differ randomly by ±1 from the corresponding parameters for their ancestor.

Probability of birth: We chose the time-dependent probability of birth per unit time as

$$p_b(i, \alpha) = \frac{X_{\text{max}}(i) - X(i, \alpha; t)}{X_{\text{max}}(i) - X_{\text{rep}}(i)} \left(1 - \frac{n_i}{n_{\text{max}}}ight)$$

if $X(i, \alpha) \geq X_{\text{rep}}(i)$ and $X_{\text{max}}(i) > X_{\text{rep}}(i)$

Note that in the limit of vanishingly small population, i.e., $n_i \to 0$, we have $p_b(i, \alpha) \to 1$ if $X(i, \alpha) = X_{\text{rep}}(i)$ and, then, $p_b$ decreases linearly as the organism grows older. However, since the eco-system can support only a maximum of $n_{\text{max}}$ individual organisms of each species, $p_b(i, \alpha; t) \to 0$ as $n_i(t) \to n_{\text{max}}$, irrespective of the age of the individual organism $\alpha$.

Probability of natural death: We assume the probability of “natural” death (due to ageing) to have the form

$$p_d(i, \alpha) = \frac{X_{\text{max}}(i) - X(i, \alpha)M(i)}{X_{\text{max}}(i)M(i) - X_{\text{rep}}(i)}$$

if $X(i, \alpha) \geq X_{\text{rep}}(i)$

$$p_d(i, \alpha) = \frac{X_{\text{rep}}(i)M(i) - X_{\text{rep}}(i)}{X_{\text{max}}(i)M(i) - X_{\text{rep}}(i)}$$

if $X(i, \alpha) < X_{\text{rep}}(i)$

provided $X_{\text{max}}(i)M(i) > X_{\text{rep}}(i)$. In all other situations, $p_d(i, \alpha) = 1$. Note that, for a given $X_{\text{max}}$ and $X_{\text{rep}}$, the larger is the $M$ the higher is the $p_d$ for any age $X$. Therefore, each species have a tendency to increase $M$ for giving birth to larger number of offsprings whereas the higher mortality for higher $M$ opposes this tendency.

The longest runs in our computer simulations were continued up to a maximum of five million time steps. If each

![FIG. 1: The total number of species $N(t)$ is plotted against time; the corresponding parameter set is: $N_{\text{max}} = 100$, $n_{\text{max}} = 1000$, $C = 0.1$, $p_{\text{sp}} = 0.001$, $p_{\text{mut}} = 0.001$.](image-url)
time step in our model is assumed to correspond to a real time of the order of one year, then the time scale of 5 million years, over which we have monitored our model eco-system, is comparable to real geological time scales.

Since we faced difficulty in getting high quality data, with reasonably good statistics, for \( N_{\text{max}} > 100 \) and \( n_{\text{max}} > 1000 \), we have carried out most of our simulations with \( N_{\text{max}} = 50, 100 \) and \( n_{\text{max}} = 100, 1000 \) only. The data obtained from the different runs, each starting from a random initial condition, were averaged. Both CRAY-T3E and SUN workstations were used for the simulations.

In fig.2 we plot the total number of species, \( N(t) \), in a particular run, starting from a single initial condition, upto half a million time steps. In fig.2 we plot the corresponding variation of the total population \( n(t) \) over relatively short interval of 20,000 time steps only. These clearly demonstrate that the evolution has periods of “stasis” during which organisms populations keep fluctuating; the stasis are interrupted by occasional bursts of rapid extinctions followed by slower recovery.

The average distributions of the lifetimes of the species are plotted in fig.3 for one set of values of the parameters. Clearly, the data are consistent with a power-law; the effective exponent, which is, approximately, 2, is also consistent with the corresponding estimate quoted in the literature [1, 2]. However, in fig.3 the power law holds only over a limited range [13]. Since real eco-systems are much more complex than our model eco-system and the available fossil data are quite sparse, it is questionable whether real extinctions follow power laws and, if so, over how many orders of magnitude.

We have also observed (not shown) that the higher is the mutation probability \( p_{\text{mut}} \), the lower is the lifetime; this is consistent with the intuitive expectation that the higher rate of mutation leads to higher levels of biological activity in the eco-system thereby leading to the extinction of larger number of species. But, \( p_{sp} \) had weaker effect on the same data. However, if \( p_{sp} \) is too small to maintain adequate pace of recovery of the eco-system after mass extinctions, the entire eco-system collapses.

Fig.4 shows the time-averaged age-distribution in the populations of a species as well as the distributions of \( X_{\text{max}}, X_{\text{rep}} \) and \( M \). We see that the minimum age of reproduction \( X_{\text{rep}} \) is quite small, as usual in the employed ageing model [1]. The age distribution decays stronger than a simple exponential, indicating a mortality increasing with age as it should be [12]. The genetic death ages \( X_{\text{max}} \) are far above the upper end \( \approx 31 \) of the age distribution, as is appropriate for animals in the wild [14]. Finally, fig.5 shows the distribution of \( M(i) \); this is relatively much broader than the distributions of \( X_{\text{max}} \) and \( X_{\text{rep}} \).

In this letter we have presented a unified model which
describes the birth, ageing and death of individuals and population dynamics on short time scales as well as the long-time evolution of species. Not only the total number of species and the inter-species interactions but also the collective characteristics, namely, \(X_{\text{rep}}, X_{\text{max}}\) and \(M\), of each species vary following a stochastic dynamics. Thus, our model is capable of self-organization. To our knowledge, there are only a few earlier evolutionary models based on inter-species interactions which describe population dynamics of each species. The population dynamics within the framework of Lotka-Volterra equations have been considered earlier for only a few species. But, these do not account for the age distributions as the entire population of each species is represented collectively by a single dynamical variable in contrast to the explicit birth, growth and death of individual organisms captured in our model.

Since we have observed strong deviations from power-law in the distributions of lifetimes of species even in the Sole-Manrubia model, in spite of the absence of detailed "micro"-dynamics in the latter, we strongly believe that this is a generic features of evolution and extinction of species. It would be interesting to investigate the geographical effects on our model eco-system by re-formulating it on a lattice in the same spirit in which some lattice models of prey-predator systems have been formulated.

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