

Competition between cryptic species explains variations in rates of lineage evolution

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Gradual evolution is a common phenomenon in the fossil record of marine microplankton, yet no theoretical model has so far been presented to explain the observed pattern of unidirectionality in trait evolution lasting over tens of millions of generations. Recent molecular genetic data show that the majority of microfossil-producing plankton groups harbors substantial cryptic diversity. Here, we examine the effect of cryptic diversity on apparent rates of lineage evolution. By using a theoretical approach, we show that under resource competition, an increasing number of sibling species within a hypothetical lineage leads to an exponential slowdown of the apparent rate of evolution. This mechanism explains both the remarkable variation in apparent rates of evolution observed in marine plankton, as well as the presence of long gradual evolutionary trends.

evolutionary rate | foraminifera | fossils | diversity | ecological dynamics

The fossil record of marine microplankton has been instrumental in providing quantitative data on the rates and patterns of morphological evolution. Contrary to terrestrial and shallow-marine settings, deep-sea sediments routinely provide long, well dated, continuous sequences documenting changes in the morphology of the fossilized remains of marine plankton at the resolution of a few thousand years (1). The data on evolutionary rates of marine microplankton (mostly planktonic foraminifera) derived from the fossil record indicate a striking range of rates of lineage evolution: whereas some transitions are completed in $<10^5$ to 10^6 generations [assuming a monthly reproductive cycle in planktonic foraminifera (2)] (3–5), long unidirectional trends in morphological traits have been documented to last $>10^7$ or even 10^8 generations (6–10) (Fig. 1).

The remarkably slow rate of morphological evolution in some marine microplankton lineages has attracted considerable attention, particularly because long-lasting gradual trends are not easy to accommodate within neo-Darwinian evolutionary mechanisms (11). However, no appropriate explanation has ever been put forward. The apparent rate of morphological evolution in these lineages appears much slower than predicted by classical evolutionary theory (12). Lande (13) postulated that the gradual patterns could represent random genetic drift, but increasingly sophisticated statistical analyses of the *Contusotrucana* lineage (10) (Fig. 1), for example, indicated a significantly directional component deviating from random null models (14, 15). An explanation involving the tracking of a gradually shifting optimum by the evolving lineages is equally illusory: on geological time scales, the variance in surface ocean properties is dominated by orbitally driven insolation changes with periods between 20 and 400 kyr, as was the case for the late Cretaceous habitat of the *Contusotrucana* lineage (16).

All earlier interpretations of gradual trends in fossil microplankton relied on the assumption that each of the evolving lineages represented a single (chrono-) species. The discovery of a prevalent cryptic genetic diversity within species of planktonic foraminifera (17) and other fossil-producing plankton (e.g., ref. 18) implies that evolutionary patterns extracted from fossil

microplankton may represent the development of a cohort of cryptic sibling species. Available data indicate that, in many cases, these sibling species can be genetically and ecologically distinct, but the morphologies of their fossilized remains cannot be distinguished (19–22). Importantly, the cryptic genetic diversity within morphologically defined species appears finite; for example, so far only two to seven genetic types have been described per morphospecies of planktonic foraminifera, with as many as four co-occurring at the same location (17). The existence of such cryptic diversity implies that the observed morphological evolution in marine microplankton lineages may need to be subdivided among the contributions of several coevolving species, whose interactions may have an impact on how, and how fast the consortium evolves.

Although the discussion of mechanisms explaining the pattern of continuous evolutionary change and stasis in fossil lineages has recently received renewed attention (e.g., 23, 24), none of the models presented to date explicitly accounts for the ecological interaction between coevolving (sibling) species. Therefore, they overlook frequency dependence processes, that is, how population dynamics are affected by the frequency of each morph in the population. Here, we investigate explicitly the consequences of cryptic diversity for the apparent rate of evolution. To do so, we formulate a theoretical model for a number of interacting morphologically cryptic species and analyze the model to determine quantitatively how cryptic diversity can affect the rate of evolution within the system. In this model, instead of an absolute fitness measure that depends only on the trait of an individual, we use a relative fitness measure that depends on the value of all of the other traits and the frequency at which they occur.

Results

Little is known about the ecological interaction between contemporary foraminifera, and even less information is available about the interactions between fossil species. Not knowing any details, we will describe the interactions with the Lotka–Volterra interaction model, which provides a general template to describe the interactions between species (25). In the absence of any details, we postulate that the interaction between foraminifera is dominated by competition for resources and formulate a model that captures the essence of competition for resources among morphologically cryptic sibling species. This chosen model is considered representative inasmuch as we expect that the general findings derived from it would be qualitatively similar to those derived from other or more complex models. We then proceed to extract the selection pressure on the different

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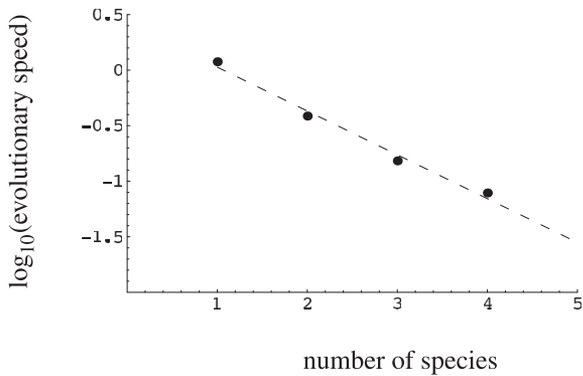


Fig. 3. Influence of the number of species on the evolutionary speed of the system. The log-linear relationship is $y = 0.42 - 0.40x$ ($R^2 = 0.987$). Parameter values are $s_{\max} = 20$ and $\gamma = 20$ (we choose this value instead of 25 to allow the coexistence of >3 species in the system).

evolutionary change. To arrive at such results we assumed that populations are at their equilibrium and that the trait variance within a species is proportional to their population size. To demonstrate that our results are robust against relaxing these assumptions, we simulated an equivalent system for which these assumptions were not made. We assumed that there is an underlying trait that is under selection (exactly as in the deterministic model) and that small mutations in this trait occasionally occur by chance. To reflect the fact that this trait is correlated, but not identical, to a morphological variable (such as the conicity of the shells) we assumed that morphological variables are normally distributed, with a constant variance and a mean that equals the value of the underlying trait. Fig. 4 shows two simulations with such a model, which we set up to accommodate either a single species or three cryptic species. The underlying evolutionary dynamics of the trait are very similar to the dynamics shown in Fig. 2 (for details, see *SI Text Appendix B*, and *Fig. S2*).

Discussion

In this study, we show how taking into account hidden diversity in an evolving lineage may help to understand patterns of evolutionary change in marine microplankton. An increase in cryptic diversity leads to an evolutionary slowdown of the evolving system through resource competition. The exact mechanism can easily be understood by considering the competition between the cryptic species (*Fig. S3*). By changing the environmental parameters, the most dominant competitor will respond first by adjusting its trait. Once this has happened, the next species in the competitive hierarchy feels the selection pressure and, consequently, the trait of this species changes. Only then can the next species in the hierarchy respond and move toward its new ecological niche. The key point to use in interpreting our results is the time factor: subdominant species can only evolve once the dominant species have evolved. For a species that has i species above it in the competitive hierarchy to evolve, i times the number of mutations are needed, and the rate of evolution of the i th species will depend on the product of the phenotypic variation of the i best competitors. This creates the observed exponential decay. This mechanism is not restricted to cryptic sibling species. However, because sibling species can be expected to compete for resources much more intensely than nonsibling species of foraminifera, the evolutionary slowdown will be more pronounced for sibling species, than similar effects resulting from competition with other organisms.

A secondary effect that explains the relatively slow evolution of assemblages of cryptic species is that these species are likely

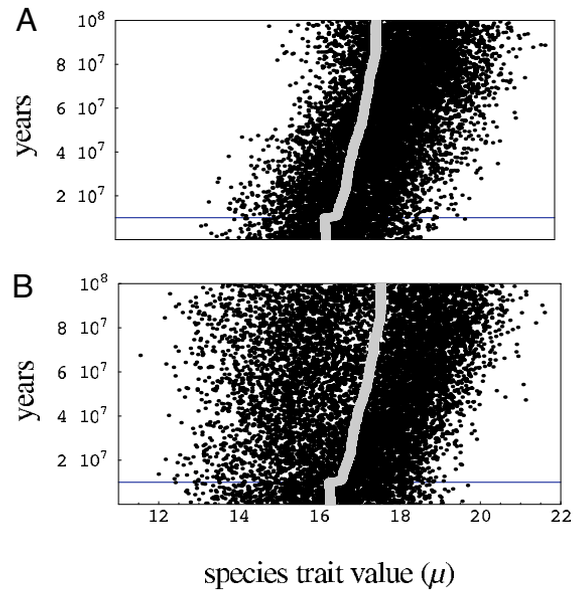


Fig. 4. Stochastic simulation of the evolution of trait μ_i in a system with one (A) or with three species (B). At $t = 10^7$ years (horizontal line), we gradually change s_{\max} from 19 to 22 (in 10^6 years). The average value of the trait (bold gray line) evolves more slowly when there are more species in the system. In both A and B, $\gamma = 25$ and $\tau = 10^{-5}$. For further details, see *SI Text Appendix B*.

to have less phenotypic variation per species, simply because the cryptic species will differ from one another. Lumping several cryptic species together into a single morphospecies will drastically overestimate the amount of variation within that morphospecies. Arguments about the rate of evolution based on the evolution of a single species thus overestimate the rate of evolution when applied to an assemblage of cryptic species.

Our model shows how and why the presence of cryptic species in an evolving lineage leads to a slowdown of apparent morphological evolution. Despite the lack of parameterization, we observe remarkable consistency between the predictions of our model and fossil data. First, analytical solutions of our theoretical model indicate that a two-orders-of-magnitude slowdown in a system with four cryptic species and a three-orders-of-magnitude slowdown could be achieved in a system with only seven cryptic species; both the implied variation in rates of evolution (*Fig. 1*) and number of cryptic species (17) agree with observations. Second, the model predicts an asymmetrical pattern in the evolution of the lineage with a more rapid change at one end of the morphospace. This feature has been observed in the *Contusotruncana* lineage (*Fig. 1*).

The phenomenon of cryptic (or sibling) species is well documented in the plankton (26), but it is by no means restricted to this ecological group, nor is there any evidence that it is more common among protists (most literature on cryptic species actually derives from insects; see, e.g., ref. 27). Therefore, in theory, our model could be applied to any group of organisms. There is, however, one major limitation: to engage in competition, the cryptic sibling species in our model must occur in sympatry and share the same resources. This often holds in the pelagic environment, which is relatively unstructured spatially. In a large compilation of fossil data, Hunt (28) shows that gradual unidirectional evolution is rare, in general, but appears more frequent among planktonic organisms. This could indicate that the mechanism described by us may be more pertinent for this environment.

It is increasingly recognized that ecological feedback can have profound effects on the evolution of traits. These insights

have been supported by both theoretical and experimental studies (29–35). Here, the application of the competition concept to a consortium of cryptic sibling species offers the first theoretical explanation for the existence of long unidirectional patterns of trait evolution observed in the fossil record of marine microplankton.

The Model

Our model has N species, where individuals of each species i are characterized by the mean size of the food particles they ingest (μ_i). This leads to N ordinary differential equations that describe the dynamics of the population density of species i denoted $x_i(t)$ and one partial differential equation to describe the change in the amount of food particles of size s denoted $n(s,t)$. The equations read:

$$\frac{\partial n(s,t)}{\partial t} = n(s,t)(1 - n(s,t)) - \sum_{j=1}^N x_j(t)k(\mu_j,s)n(s,t) \quad [1]$$

$$\frac{dx_i(t)}{dt} = x_i(t) \int_0^{s_{\max}} k(\mu_i,s)n(s,t)c(s)ds - \gamma x_i(t), \quad [2]$$

where γ is the mortality rate of all species, $c(s)$ is the energetic value of food particles of size s , and s_{\max} is the size of the largest particles. The kernel function $k(\mu_i,s)$ describes the proportion of food particles of size s eaten by a species that ingests, on average, food particles of size μ_i .

To analyze the above system, we assume a homogeneous carrying capacity for the food particles population such that, without consumption, at equilibrium, Eq. 1 becomes $\tilde{n}_0(s) = 1$. (Here, as in the following, tildes indicate equilibrium values.) Next, as in previous studies (36), we choose a kernel function that has a double-exponential shape centered around μ_i (SI Text Appendix A, Choice of a kernel function). Finally, we choose an energetic-value function that depends linearly on the food particle's size, that is, $c(s) = s$ (taking the energetic value proportional to volume would not modify the results qualitatively and would complicate the calculations).

If the resource population reaches its equilibrium much faster than the plankton populations [i.e., that $n(s,t) \approx \tilde{n}(s)$], we can easily reformulate our equations to a classical Lotka–Volterra competition model (SI Text Appendix A, Derivation of the Lotka–Volterra model):

$$\frac{dx_i}{dt} = x_i \left(R(\mu_i) - \sum_{j=1}^N x_j(t)\alpha(\mu_i, \mu_j) \right), \quad [3]$$

where $R(\mu_i) = \int_0^{s_{\max}} k(\mu_i,s)c(s)ds - \gamma$ and $\alpha(\mu_i, \mu_j) = \int_0^{s_{\max}} k(\mu_i,s)k(\mu_j,s)c(s)ds$. As we show in SI Text Appendix A, Finding the equilibrium densities of the populations, obtaining the species equilibrium densities (\bar{x}_i) from Eq. 3 is straightforward. These equilibrium densities depend on $R(\mu_i)$ and $\alpha(\mu_i, \mu_j)$.

We then use Eq. 3 to derive the invasion fitness function of a rare mutant. The mean particle food size of a mutant (μ_i^*) differs slightly from that of the resident (μ_i). After some simplification

(SI Text Appendix A, Introducing a mutant), the equation describing a mutant's density can be written as

$$\frac{dx_i^*}{dt} \approx \left(R(\mu_i^*) - \sum_{j=1}^N x_j\alpha(\mu_i^*, \mu_j) \right) x_i^* \quad [4]$$

The invasion fitness W_N of a mutant of resident species i in a system with N species is given by

$$W_N(\mu_1, \dots, \mu_i, \dots, \mu_N, \mu_i^*) = \frac{1}{x_i^*} \frac{dx_i^*}{dt} \quad [5]$$

From Eq. 4, we get

$$W_N(\mu_1, \dots, \mu_i, \dots, \mu_N, \mu_i^*) = R(\mu_i^*) - \sum_{j=1}^N \bar{x}_j\alpha(\mu_i^*, \mu_j). \quad [6]$$

The fact that the mutant is rare (compared with the resident) allows us to assume that resident species do not “feel” the presence of a mutant population. If the resident populations are also at equilibrium, the density values x_j can be replaced by their equilibrium values \bar{x}_j .

From Eq. 6, we derive the effect on the invasion fitness of a mutation causing a small change in a resident trait μ_i . This marginal fitness gives us the value of the selection differential (sd) of a mutant of resident species i , which indicates in which direction the trait evolves and at which speed. The selection differential value is obtained by deriving $W_{i,N}$ with respect to μ_i^* ,

$$sd_{i,N} = \left. \frac{dW_N(\mu_1, \dots, \mu_i, \dots, \mu_N, \mu_i^*)}{d\mu_i^*} \right|_{\mu_i^*=\mu_i} \quad [7]$$

See SI Text Appendix A, Calculating the selection differential and the evolutionary rate, and Figs. S4 and S5, for further details. If $sd_{i,N} = 0$, species i is at an evolutionary equilibrium. If $sd_{i,N} > 0$, mutants with $\mu_i^* > \mu_i$ will be selected for and if $sd_{i,N} < 0$ mutants with $\mu_i^* < \mu_i$ will be selected for. Because we have a community of N species, which may all create mutants, we get N values for the selection differential of the system.

Knowing the selection differentials allows us to quantify the rate of the evolutionary process and follow variations in trait value (μ_i). Generally, the rate of evolution is the product of the selection differential and the amount of heritable variation (13, 37, 38). The amount of variation generally will scale with number of mutations, which is proportional to the size of the population and the mutation rate (38). Therefore, we can write the rate of evolution in the mean food particle size of any species i in the system as

$$\frac{d\mu_i(t)}{dt} = \tau \bar{x}_i sd_{i,N}, \quad [8]$$

where τ is a constant proportional to the mutation rate, which is the same for all species in the community.

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