Extant mammal body masses suggest punctuated equilibrium

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1. INTRODUCTION

Charles Darwin apparently hypothesized not only that natural selection causes microevolutionary changes in the average phenotype of populations from one generation to another, but also that, accumulated over thousands of years, these same changes account for phenotypic differences between species (Mayr 1982; Gould 2002).

In the fossil record, however, species often appear suddenly different, to remain largely unchanged until they disappear again (Stanley 1998). For example, the European mammoth survived the climatic vicissitudes of the last ice age with apparently little phenotypic change, as Hugh Falconer pointed out to Darwin, after which Darwin added to the third edition of ‘The origin’ the following: ‘It is a more important consideration, clearly leading to the same result, as lately insisted on by Dr. Falconer, namely, that the periods during which species have been undergoing modification, though very long as measured by years, have probably been short in comparison with the periods during which these same species remained without undergoing any change’ (Stanley 1981, 1998).

Paradoxically however, after the success of the theory of microevolution by means of natural selection in the twentieth century, Darwin’s idea of macroevolution as the long-term consequence of microevolution was the ruling paradigm when, just over 35 years ago, Eldredge & Gould (1972) introduced their theory of ‘punctuated equilibrium’. This theory, which hypothesizes that interspecific morphological change is concentrated in rapid bursts triggered by speciation, has fuelled speculation about the mode of character evolution ever since (Gould & Eldredge 1993; Gould 2002). The theory proved hard to test, however, because speciation typically occurs on a time span many times the duration of a biological experiment, and because species from fossil remains can be distinguished only if morphologically distinct (Stebbins & Ayala 1981). Cryptic species, for example, would not be distinguished in the fossil record, morphologically similar ecomorphs (Losos et al. 1998) would go unnoticed, and replacement of local ecotypes with populations from elsewhere might be mistaken as a sudden morphological shift. Consequently, the importance of gradual and ‘punctuational’ evolution for the origin of morphological diversity in biological species remains largely unknown.

Fortunately, fossil morphology is not the only source of information concerning the mode of evolution. Molecular phylogenies substitute for the time frame provided by the fossil record and allow gradual evolution and punctuated equilibrium to be distinguished based on phenotypes of extant species (Pagel 1999; Pagel et al. 2006): if phenotypic evolution is concentrated in short periods of time when new species emerge, even young sister species could show significant phenotypic differentiation, whereas Darwin’s original view of gradual change implies that interspecific phenotypic variance requires sufficient time to develop (Avise 1977). Therefore, we can estimate separately the rates of both speciational and gradual phenotypic evolution from a phylogeny and phenotypes of present-day species, without reference to fossil morphology (Avise 1977; Ricklefs 1980; Douglas & Avise 1982; Lemen & Freeman 1989; Mooers & Schluter 1998; Bokma 2002).

We estimated rates of gradual (anagenetic) and speciational (cladogenetic) body size evolution from body masses of 2143 of 4510 present-day mammal species on a composite phylogenetic tree that represents 99% of existing species (Bininda-Emonds et al. 2007). A novel Bayesian algorithm (Bokma in press) allowed us to simultaneously estimate speciation and extinction


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probabilities, thus taking into account uncertainty in the number of extinct lineages and speciation events that do not show on a phylogeny reconstructed from extant species. Thereby, we overcome the major drawback of previous methods to estimate the mode of evolution from neontological data (Avise 1977; Ricklefs 1980; Douglas & Avise 1982; Lemen & Freeman 1989; Mooers & Schluter 1998).

2. MATERIAL AND METHODS

(a) Estimation algorithm

We estimated rates of anagenetic and cladogenetic body size evolution assuming a branching Brownian motion model of evolution of the logarithmic phenotype (Bokma 2002), with speciation and extinction rates \( \lambda \) and \( \mu \) and variances of Brownian motion \( \sigma^2 \) per Myr (anagenetic) and \( \sigma^2 \) at the instant a new species is born (cladogenetic). \( \sigma^2 = 0 \) means evolution is purely gradual; that is, newborn species have the same phenotype as their parent. According to this model, the logarithmic body size change over a branch of the phylogenetic tree is normally distributed with zero mean and variance \( \mu b_{o} \sigma^2 + S_o \sigma^2 \), where \( b_o \) is the branch length, \( S \) is the number of speciations on the branch, and \( \sigma^2 \) and \( \sigma^2 \) are the ‘rates’ of evolution per unit time and speciation event, respectively (Bokma 2002). It may appear counter-intuitive that a branch contains speciation events, but we are analysing a reconstructed phylogeny (since Nee et al. 1994), pruned of extinct lineages. The speciation events that do not appear on the reconstructed tree may have affected the phenotypes of extant species (Bokma 2002). Given \( b_o \) and \( S \), the likelihood of body size change over a branch depends only on \( \sigma^2 \) and \( \sigma^2 \) and the likelihood of the tree is simply the sum of its branches’ likelihoods.

Obviously, \( S \) is not known, and neither are the body size changes over the branches, because we only have body sizes of some existing species. Therefore, all unknown parameters (i.e., \( S \), the body sizes of all ancestral species, and the body sizes of extant species for which we had no measurements) were estimated from Monte Carlo Markov chains constructed by Metropolis–Hastings sampling, started from arbitrary initial values (Bokma in press). Inference of \( S \) is the critical part of the estimation procedure. \( S \) consists of two parts: \( S = S_o + S_h \), where \( S_o \) is the speciation event that marks the origin of the branch (and can be observed from the tree) and \( S_h \) is the number of speciations that are hidden on the branch. We assume that only newborn species undergo cladogenetic evolution, so that \( S_o \) is either 1 or 0, depending on whether or not the focal branch represents the newborn species. Of course, we cannot detect from the tree alone which was the newborn species, but the phenotypic data may favour one assignment over the other. \( S_h \) may assume any (integer) value. Given probabilities of speciation and extinction \( \lambda \) and \( \mu \), respectively, and assuming \( \lambda \) and \( \mu \) are constant across the tree, \( S \) can to a good approximation be sampled from a Poisson distribution (Bokma in press), \( \lambda \) and \( \mu \) in turn can be estimated from the phylogenetic tree (Nee et al. 1994; Bokma 2003). Here, we used the likelihood of the distribution of species over the 83 taxa older than (arbitrarily) 50 Myr defined by the phylogenetic tree (Bininda-Emonds et al. 2007) to Metropolis–Hasting sample \( \lambda \) and \( \mu \). When analysing sub-clades (see below), we estimated \( \lambda \) and \( \mu \) from the branching times of the sub-clade tree with a Bayesian modification of Nee et al.’s (1994) method that allowed us to simultaneously estimate uncertain branching times resulting from polytomies.

The composite phylogeny used for the present analyses (Bininda-Emonds et al. 2007) contains many polytomies: 27% of all branches. Because we cannot evaluate gradual phenotypic change on zero-length branches, and because the lengths of these branches are obviously incorrect, zero-length branches as well as their immediate descendant branches were excluded from all analyses (except estimation of \( \gamma \) and \( \mu \)). Estimates were obtained from every 50th sample from MCMC chains of 50 000 iterations after a burn-in of 5000 iterations.

(b) Data

Body size data (expressed in kg or ln(kg)) throughout were obtained mainly from Silva & Downing (1995). Where multiple measurements were reported per species, mean, minimum and maximum body sizes were averaged, if applicable per sex. If only a range of body sizes was available, the average was calculated as the range midpoint. Subsequently, if available the average of both sexes was used, otherwise the average of whatever data available. These average body masses are only estimates of the actual mean body size of a species, and therefore the latter was sampled during analyses from a normal prior distribution centred on the logarithm of the calculated average. If data from both sexes were available, the prior was assigned standard deviation 0.025, otherwise 0.05 (logarithmic scale). Considering an average intraspecific phenotypic variance of 0.14m, where \( m \) is species average body size (see below) and assuming \( m \) were calculated from approximately 10 individuals per species, our priors are rather wide. (For comparison, the interspecific standard deviation of logarithmic body masses of all mammals is 3.10).

Assuming that the minimum and maximum body sizes reported in the literature typically enclose 95% of individuals, we calculated the phenotypic standard deviation \( s \) as a quarter of the reported size range. Regression of \( s \) on \( m \) yielded a neat log–linear relationship \( \ln(s) = \ln(m) - 2 \), implying that \( s = 0.14m \).

3. RESULTS

(a) Empirical estimates

First, we set \( \sigma^2 = 0 \) and estimated \( \sigma^2 = 0.0696 \) (95% CI 0.0631–0.0778) from the complete phylogenetic and body mass data. This is the estimated variance of logarithmic body mass change over a Myr if species do not change upon speciation. It is a useful reference value, as it does not depend on \( \lambda \), \( \mu \) or \( S_h \). The rate of gradual evolution \( \sigma^2 \) refers to evolution on a logarithmic scale, but it has a straightforward interpretation on the ‘conventional’ body mass scale (Lynch 1990; Bokma 2002): over a Myr, the expected body mass change of a mammal species, expressed as a fraction of its mean body mass is \( \sigma^2 \times 0.0696 \times 0.26 \). Subsequently, we released \( \sigma^2 \). Estimating both rates of evolution simultaneously, we obtained the significantly lower estimate \( \sigma^2 = 0.0024 \) (0.0002–0.0133) and \( \sigma^2 = 0.164 \) (0.135–0.192). Now also \( \sigma^2 \) is affected, through \( \sigma^2 \) by the estimates of \( \lambda \), \( \mu \) and \( S_h \), which explains the larger confidence intervals (figure 1).

The above calculations assume that body mass changes in all mammal species have come from the same two normal distributions with variances \( \sigma^2 \) and \( \sigma^2 \), but it is possible that different parts of the phylogenetic tree have
In the absence of natural selection, the per-generation change in the average phenotype of a population depends only on its genetic variation and population size, in such a way that $s^2_\text{obs} = h^2 s^2 / (2N)$ (Lande 1976), where $h^2$ is heritability, $N$ is effective population size, $t$ generation time, and the phenotypic standard deviation $s = 0.14$ (see §2). A conservative estimate is obtained assuming a relatively high long-term effective population size $N = 10^6$ and a relatively low heritability $h^2 = 0.3$ for body size, with for mammals an approximately average generation time of 2 years; $s^2_\text{obs} = 0.29$. Even this conservative estimate is substantially higher than most of the present and most earlier (Lynch 1990) estimates.

In the absence of selection and mutation, the expected phenotypic divergence of a newborn species equals twice the additive genetic variance of the original species (Lynch & Hill 1986), or $2h^2 s^2$. Substituting $h^2 = 0.3$ and $s = 0.14$, we obtain an approximate neutral expectation of $s^2_\text{obs} = 0.11$. This should be regarded as a minimum expectation, since the heritability of body mass may well be higher than 0.3, and because genetic variance due to mutation may cause higher rates of neutral speciational evolution. This is a neutral expectation, and selection may drive speciational evolution at considerably higher rates, yet even small populations under weak selection may reach their new optimum phenotype in only a few thousand or even hundred generations (Lynch & Hill 1986).

4. DISCUSSION

A crucial assumption in the present analysis is that $\lambda$ and $\mu$ as well as $s^2_\text{obs}$ and $s^2_\text{grad}$ have been equal everywhere on the phylogenetic tree. Analysis of sub-clades indicates that this assumption may be violated: some relatively closely related species show truly large differences in body size and sub-trees including such species lead to high estimates of rates of evolution with wide confidence intervals (table 1). There are two simple and non-exclusive explanations hereof: directional selection on body size or frequent speciation (and thus frequent cladogenetic evolution) in a few lineages. It is difficult to distinguish between these two scenarios, but differences in rates of evolution—whether true differences in $s^2_\text{obs}$ and $s^2_\text{grad}$ or caused by differences in $\lambda$ and $\mu$—may lead to incorrect estimates, with for example $s^2_\text{grad}$ explaining more than 100% of observed interspecific body size variation. Thus, the clearly aberrant analyses summarized in table 1 as well that of all mammals, including all aberrant lineages, should be regarded with caution. Clades without apparently unusual (observed) size differences in closely related species all suggest gradual evolution is responsible for 20–35% of interspecific variation.

The theory of punctuated equilibrium has often been discussed with respect to the mode of speciation, perhaps because it was originally presented as the geological consequence of Mayr’s theory of allopatric speciation (Eldredge & Gould 1972). We assumed that only newborn species undergo cladogenetic evolution, which has often been linked to the idea of peripatric species formation, where punctuational change takes place when new species arise from small peripherally isolated populations. If, on the other hand, punctuational change results from divergent selection in young sister species in (secondary) sympatry, a model in which both sister species undergo cladogenetic evolution might be more
appropriate. However, as has repeatedly been pointed out (Gould 2002), several modes of speciation could as well result in rapid morphological change on a geological time scale, and the present results should not be interpreted as evidence for a particular mode of speciation.

The main significance of the present results concerns another controversy associated with punctuated equilibrium: to what extent macroevolution is the long-term consequence of microevolution and to what extent it represents fundamentally different processes. Above we have shown that even if \( s^2 = 0 \), estimated rates of gradual evolution are much higher than expected under neutral mutation–drift equilibrium, which is in line with previous observations (Lynch 1990). Stabilizing selection could be, but certainly does not have to be (Hansen & Houle 2004), the explanation why observed rates of gradual evolution are an order of magnitude lower than genetic variation would allow for.

Similarly, we showed that both the speed and magnitude of speciational evolution, even though the latter is perhaps overestimated here, could be obtained with only weak selection. Thus, the observation that the rate of phenotypic (and genotypic (Pagel et al. 2006)) evolution is elevated during speciation is not at odds with Darwin’s idea of the resulting phenotypes being the products of natural selection, and provides insufficient evidence that micro- and macroevolution would be decoupled processes. Nevertheless, explaining perhaps one-third of interspecific variation, gradual evolution seems grossly overvalued.

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