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Primate extinction risk and historical patterns of speciation and extinction in relation to body mass

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Body mass is thought to influence diversification rates, but previous studies have produced ambiguous results. We investigated patterns of diversification across 100 trees obtained from a new Bayesian inference of primate phylogeny that sampled trees in proportion to their posterior probabilities. First, we used simulations to assess the validity of previous studies that used linear models to investigate the links between IUCN Red List status and body mass. These analyses support the use of linear models for ordinal ranked data on threat status, and phylogenetic generalized linear models revealed a significant positive correlation between current extinction risk and body mass across our tree block. We then investigated historical patterns of speciation and extinction rates using a recently developed maximum likelihood method. Specifically, we predicted that body mass correlates positively with extinction rate because larger-bodied organisms reproduce more slowly, and body mass correlates negatively with speciation rate because smaller-bodied organisms are better able to partition niche space. We failed to find evidence that extinction rates covary with body mass across primate phylogeny. Similarly, speciation rate was generally unrelated to body mass, except in some tests that indicated an increase in speciation rate with increasing body mass. Importantly, we discovered that our data violated a key assumption of sample randomness with respect to body mass. After correcting for this bias, we found no association between diversification rates and mass.

**Keywords:** Primate phylogeny; speciation; extinction; IUCN Red List; body mass; diversification; Bayesian phylogenetics
1. INTRODUCTION

Several studies have implicated body size as a risk factor for extinction in mammals (Purvis et al. 2000; Cardillo et al. 2005; Clauset & Erwin 2008; Davies et al. 2008). Large body size is thought to correlate with higher extinction risk through a number of life-history covariates of increased body mass, such as longer generation times and smaller litters (Cardillo et al. 2005; Isaac et al. 2005). These factors should increase the time needed to recover from stochastic demographic reductions in population size, thus increasing the probability of extinction. To test this hypothesis, studies have quantified extinction risk among living species as a ranked variable derived from the IUCN Red List conservation status categories (www.iucnredlist.org, Purvis et al. 2000; Cardillo et al. 2005; Davies et al. 2008). In mammals, Cardillo et al. (2005) found that “intrinsic” biological life history traits influenced extinction risk only when species were >3 kg in mass. Below 3 kg mass, the primary determinants of extinction risk were “extrinsic” factors that are not genetically heritable, such as geographic range size and nearby human population density (Cardillo et al. 2005). Studies also have failed to detect a consistent association between diversification rate and body mass across mammalian clades (Gittleman & Purvis 2000; Isaac et al. 2005).

Understanding the predictors of present day extinction risk, as reflected by the IUCN Red List, has clear relevance given the pressing need for primate conservation (Cowlishaw & Dunbar 2000). For example, the great apes are all large-bodied and critically endangered (IUCN Red List). An important question is whether large bodied primate species experience higher extinction rates irrespective of human activities. If so, then large bodied primates may be especially vulnerable to anthropogenic drivers of
extinction, such as reductions in habitat or climate change. It is possible to infer
speciation and extinction rates from a dated phylogeny because higher rates of extinction
relative to speciation should produce longer internal branches on a tree with an apparent
burst of diversification close to the tips (Nee et al. 1994; Nee 2001). Recent advances
provide a way to integrate the study of speciation and extinction that is dependent upon
another trait (Maddison et al. 2007), such as body mass. Previous studies have
investigated patterns of diversification more generally in primates (Purvis et al. 1995;
Gittleman & Purvis 1998; Chan & Moore 2002; Moore et al. 2004; Paradis 2005;

Using a newly inferred primate phylogeny that enabled us to incorporate
phylogenetic uncertainty, we investigated the links between body mass, extinction risk
and diversification rates. First, we tested for an association between body mass and
present-day extinction risk categories obtained from the IUCN Red List. As with
previous studies, we predicted that larger-bodied primates are at greater risk of extinction
(Purvis et al. 2000; Cardillo et al. 2005; Davies et al. 2008). In our analyses, we used
methods that incorporate phylogenetic uncertainty, that better model character evolution
on the tree, and that test for spurious results arising from the ordinal (rather than integer)
measurement of extinction threat categories.

Second, we predicted that primate lineages characterized by greater body mass
have experienced higher extinction rates throughout evolutionary history. For this, we
applied a new method to estimate extinction rates on a phylogenetic tree in relation to a
biological characteristic (Maddison et al. 2007), again running the test in a way that
incorporated phylogenetic uncertainty.
Lastly, we turned our attention to speciation by predicting that lineages with smaller body mass have higher speciation rates. We made this prediction based on several studies showing a trend for smaller bodied mammalian clades to be more diverse, possibly because they experience vicariance events more frequently or they are able to partition the environment into more niches (Maurer et al. 1992; Gittleman & Purvis 1998; Gardezi & da Silva 1999; Purvis et al. 2000; Cardillo et al. 2005; Isaac et al. 2005).

2. METHODS

(a) Tree Inference

As our hypotheses pertained to the order Primates as a whole, we needed a phylogeny that included as many species as possible. To this end, we used the trees available from the 10kTrees project, Version 1 (http://10ktrees.fas.harvard.edu/). The website provides extensive documentation on the tree inference, a graphical interface for downloading trees, and a number of visualizations of the trees. Details regarding tree inference are available in Arnold et al. (2010). Our use of Bayesian tree inference enabled us to deal with phylogenetic uncertainty by running comparative tests on multiple trees saved from the Markov chain. Allowing for such topological and branch length uncertainty is important because the phylogeny used can affect the conclusions that are drawn from a comparative analysis (Lutzoni et al. 2001, Pagel & Lutzoni 2002).

We used two sets of trees from the Bayesian analysis: a sample of 100 trees distributed evenly along the post-burnin Markov chain and a consensus tree of all nodes with clade credibility support greater than 0.5. We dated all trees prior to comparative analysis by using seven fossil calibration points employed by previous phylogenetic
studies (Table 1, Seiffert et al. 2003; Yang & Yoder 2003; Yoder & Yang 2004; Godinot 2006; Hodgson et al. 2009). We conducted molecular dating with the software r8s (Sanderson 2002) using the penalized likelihood algorithm with a smoothing parameter of 100, chosen because this value best recovered dates inferred from phylogenetic analyses of smaller taxonomic samples but with more extensive sequence data (Yang & Yoder 2003; Yoder & Yang 2004; Hodgson et al. 2009).

(b) Body mass and IUCN Red List data

Body mass data were obtained from Smith & Jungers (1997). We calculated the mean female body mass across study sites for our analysis. In this manner, we obtained body mass data on 160 species that could be matched to our phylogeny through translation via the taxonomy of Wilson & Reader (2005). Analyses used the natural log of female body mass. We obtained IUCN conservation status for the species on our phylogeny from the IUCN Red List website (www.iucnredlist.org). From each IUCN category we constructed an ordinal variable with higher ranks corresponding to greater extinction threat. Specifically, ranked from smallest to highest, we used the following categories of threat: least concern, near threatened, vulnerable, endangered and critically endangered. Six species labeled “data deficient” in the IUCN Red List were removed from the analysis, reducing the sample size to 154 species. The highest rank was assigned to “critically endangered” rather than “extinct in the wild” because our study included only extant species as data points.
(c) Comparative analyses

We tested for an association between body size and IUCN extinction risk through a phylogenetic generalized least squares (PGLS) model applied across our Bayesian tree block. This analysis of the primate data replicates previous independent contrasts analyses across mammals (e.g. Cardillo et al. 2005; Davies et al. 2008). However, PGLS implemented in the program BayesTraits (Pagel & Meade 2007) enabled us to incorporate Bayesian estimation of the branch scaling parameter $\lambda$ (Pagel 1999) across a tree block, which should improve upon contrasts-based approaches. In particular, we suspected that Brownian motion along the branch lengths in our block of 100 trees might not reflect evolutionary change in a character such as threat status. Brownian motion along branch lengths is an assumption of independent contrasts. The scaling parameter lambda adjusts the internal branch lengths with a length multiplier such that the data meet the assumption of Brownian motion. By systematically searching through many possible values for lambda, the program locates the value that makes the data most likely and thus best accommodates the assumption of Brownian motion. In primates, Purvis et al. (2005) showed that IUCN rank alone exhibited a lambda equal to 0.77 that excluded both zero and one (95% CI 0.51-0.90). Thus, we considered it important to incorporate lambda in our PGLS, and especially to do so in a way that incorporates uncertainty in phylogenetic relationships and branch lengths.

The use of PGLS or independent contrasts treats IUCN extinction risk as a continuous variable (Felsenstein 1985; Garland et al. 1992). Counter to this assumption, extinction risk codes in the IUCN Red List are not continuously varying. Instead, extinction risk is an ordinal variable in which ranks likely vary in the amount of
difference in the actual underlying extinction risk (Purvis et al. 2005). For example, the 
true (quantitative) difference in extinction risk may differ between categories of near-
threatened and vulnerable, as compared to endangered and critically endangered. 
Treating ordinal variables as continuous can produce elevated Type 1 error rates because 
such treatment applies arithmetic operations that do not preserve the variance structure of 
the original ordinal ranks (Stevens 1946; Siegel 1956). The problem occurs specifically 
when the ordinal ranks are separated by unequal distances along the underlying 
continuous variable that they measure, a point acknowledged by Purvis et al. (2005) and 
given in the example above.

We used computer simulations to assess whether the treatment of IUCN threat 
categories as continuously varying may have introduced error into this study and previous 
 studies (Cardillo et al. 2005; Purvis et al. 2005; Davies et al. 2008). Specifically, we 
tested whether treating threat status as continuously varying results in elevated Type 1 
error rates. We conducted 1000 simulations of two uncorrelated continuous characters on 
Characters evolved randomly with a constant accumulation of variance (set at 1.0 per unit 
branch length) and an initial state of zero. We then rescored one character from each pair 
into a set of ordinal ranks with the same number of species in each state as was observed 
for each corresponding IUCN threat status in our data set. The lowest continuous values 
of the simulated character thus became rank “0,” while the highest became rank “4,” with 
other ranks derived from intermediate values and all in matching proportion to the 
observed frequency of each rank in the IUCN data. This rendered one character a true 
continuous distribution, like body size, but the other character had been rescored into a
set of ordinal ranks with different real distances between the means of each rank. We then conducted 1000 PGLS tests to test for a significant association of the two characters, one of which was ordinal and one continuous. Significant associations were counted as Type 1 errors because the characters evolved independently during the simulations. An alternative approach to deal with ordinal IUCN data would be to include additional parameters that model the ordinal nature of the dependant variable. While this procedure in principle can be implemented (Hadfield & Nakagawa 2010), given the prominent prior research that treated IUCN as continuous, we prefer to use simulations to test this approach more generally.

To test for effects of body size on the actual speciation and extinction rates experienced by different lineages on the primate tree, we employed the “binary-state speciation and extinction” (BiSSE) test (Maddison et al. 2007), as implemented in the R package diversitree (FitzJohn 2009; FitzJohn et al. 2009). This procedure uses likelihood methods to test a six-parameter model of speciation, extinction, and trait evolution. Following the procedure recommended in Maddison et al. (2007), we constructed five parameter and six parameter models to test body-size dependant speciation and extinction rates, which we investigated separately. For each state of a binary character, BiSSE can model a rate of speciation, extinction, and character state transitions. Thus, six rates are possible for the most complex BiSSE model. The five-parameter models constrained the speciation (or extinction) rates to be equal in the two body size categories, while the six-parameter models allowed speciation (or extinction) rates to vary for different body sizes. Because these models were nested, we assessed statistical significance using likelihood ratio tests. We conducted all BiSSE analyses on our dated consensus tree.
To use BiSSE with our measures of body mass, we binned species into categories of “small” and “large” body mass. As it is not immediately clear what cutoff should be used for these categories, we initially tested our hypotheses with three different cutoffs derived from the primatological literature. The first break point, 500 g, reflects the point above which primate species are not strictly faunivorous (i.e., Kay’s threshold, Kay 1984). We thought this widely accepted energetic constraint might reflect the life history variables that underlie previously observed associations between body mass and extinction risk. The second break point, 984 g, was the phylogenetic mean body mass at the root of the dated consensus tree, as inferred through squared change parsimony reconstruction (Maddison 1991) of log body mass values in Mesquite version 2.6 (Maddison & Maddison 2006). The third break point of 3000 g came from a study of extinction risk across mammals (Cardillo et al. 2005), in which the authors found that biologically intrinsic life history traits only influenced extinction risk above a body mass of 3000 g. We ran these analyses of the three a priori break points across the 100 dated trees from the Bayesian tree search.

To assess the sensitivity of our results to the break point used for binning species, we conducted an analysis with a sliding break point that divided the body mass data into small and large categories along 20 intervals of 0.2 log units of body mass. We quantified the precision of the sliding break point estimates by sampling the six-parameter model with a Monte-Carlo Markov Chain (MCMC) that we initialized with the maximum likelihood estimates for parameter values (R package diversitree, FitzJohn 2009; FitzJohn et al. 2009). We used only the dated consensus tree for the sliding break point analysis.
The implementation of BiSSE in diversitree allows one to specify the degree of species sampling employed in the study. Doing so is important because methods to test trait-dependent extinction are biased by incomplete sampling of the study group (Nee et al. 1994; FitzJohn et al. 2009). The diversitree package includes modified likelihood equations to account for this bias (FitzJohn et al. 2009). Under the Wilson and Reeder (2005) taxonomy, which served as the taxonomy for the present study, we sampled 42% of extant Primates. We used this value as our sampling percentage in all BiSSE analyses.

The BiSSE correction for incomplete sampling is statistically valid only when the species have been sampled randomly. Two issues are relevant here: random sampling of species from the phylogeny, and random sampling with respect to the character hypothesized to affect speciation and extinction rates.

To assess the randomness of our species sample with respect to phylogeny, we conducted a G-test of proportions of species from each genera within our observed sample (Sokal & Rohlf 1995; R script). The G-test uses maximum likelihood techniques to assess whether an observed proportion of species within genera can be viewed as a random sample of species given their known frequencies from the complete taxonomy (Wilson & Reeder 2005). We conducted this test at the generic level because, based on our primate phylogeny (Arnold et al. 2010), generic classifications reflect monophyletic clades.

To test whether sampling was random with respect to body mass, we calculated the deviation of the observed number of species sampled from a genus compared to the number expected to be sampled given the number of species in each genus in the complete taxonomy. Thus, a positive deviation indicated that more species were sampled
from a genus than expected under random sampling, while a negative number indicated
fewer were sampled than expected. We then conducted a phylogenetic generalized least
squares (PGLS) test for an association between body mass and sampling deviation using
our dated consensus tree to control for phylogenetic non-independence.

We then retested our hypothesis regarding speciation and body size using the
diversification test of Freckleton et al. (2008). This test is simply a PGLS of a character
trait as a dependant variable with the number of nodes from the tree root to each tip as the
independent variable. If a particular character state, such as body size, is associated
positively with diversification rates, then a positive association should exist between the
character value and lineages that exhibit more nodes. The advantage of this test is that,
because it is conducted through a PGLS framework, we were able to include the
deviations from the randomly expected sampling as an independent variable; in other
words, we could investigate the effects of body mass on diversification while controlling
for body mass-related biased sampling if it exists, even if not statistically significant in
the above tests. Another advantage of this test is it treats body size as a continuous
variable, whereas the use of BiSSE required us to bin body size into a set of binary
categories. The disadvantage of the Freckleton et al. (2008) test is that it investigates the
association between net diversification rate and body size, rather than specifically
investigating speciation and extinction separately (as in BiSSE). We used the dated
consensus tree for this analysis.

3. RESULTS

(a) Extinction risk and body mass
In our dataset, 68 species were categorized as “Least Concern”, 9 as “Near Threatened”, 30 as “Vulnerable”, 40 as “Endangered”, and 7 as “Critically Endangered” (Supplemental Table 1). We found a significant and positive association between IUCN extinction risk and body mass while in a PGLS analysis across our dated Bayesian tree block (N = 154, \( \beta = 0.40, p = 0.004 \)). Additionally, we found that lambda differed from zero and one (\( \lambda = 0.78 \), 95% credible interval \( \lambda = 0.63 – 0.89 \)). This indicates significant phylogenetic signal and thus highlights the importance of controlling for phylogeny, but further shows that a non-Brownian component may contribute to variation in residual values. In our 1000 simulations that tested the effect of ordinal ranking for IUCN, only 5% of statistical tests were deemed significant at the 0.05 nominal level.

**Extinction rate and body mass**

In our BiSSE analyses of 160 primate species, likelihood ratio tests supported no significant relationship between log body mass and historical extinction rate either at our \textit{a priori} break points or in our sliding break point analysis (Figure 1a). Similarly, across the 100 trees, extinction rates at the \textit{a priori} break points were significantly different relative to body mass for only 1 of the 100 trees sampled from our tree block. This was true even though extinction rates were estimated to be different from zero in many cases (Figure 1a). Several papers have indicated that methods for inferring trait dependant extinction may suffer from low statistical power (Maddison \textit{et al.} 2007; Kubo & Iwasa 1995; Purvis \textit{et al.} 1995; Gardezi & da Silva 1999; Paradis 2004), a point to which we return in the Discussion.
(c) Speciation rate and body mass

In contrast to the results from the extinction analysis, we found significant speciation rate differences at the Kay’s threshold *a priori* break point on our dated consensus phylogeny and across 95 of the 100 trees (for the consensus tree, \( p = 0.016 \)). Differences in speciation rate at the phylogenetic mean and 3 kg break points were not significant on the consensus tree or on any of the trees from the tree block. Importantly, the result for Kay’s Threshold was opposite to our prediction, with higher estimated speciation rates in larger-bodied species (Figure 1b). This pattern seemed to apply across most of the variation in primate body mass, but not at the largest body mass categories (Figure 1b).

However, MCMC sampling of the speciation estimates showed that 95% credible intervals overlapped across most of the distribution (Figure 1b).

In testing for speciation and extinction rates, BiSSE also estimated two rates of transition (small to large; large to small) in all analyses. The transition rate estimates showed overlapping credible intervals across almost the entire range of sliding break points (Supplemental Figure 1). This indicates rates of body size increase and decrease are generally equivalent across the range of primate body sizes.

The BiSSE results are not compromised by nonrandom sampling of species in our dataset with respect to phylogeny. The G-test for independence showed that the observed sampling of species within genera was within the variation expected from random sampling of taxa (\( p = 0.147 \)). However, the observed sample of species was biased toward greater sampling of species with larger body mass in a PGLS test (\( \beta = 0.80, p = 0.037 \)). Thus, biased sampling towards species with larger body mass may have influenced the BiSSE results; specifically, the denser sampling of the larger species may
have biased the test to estimate a higher rate of speciation because these lineages would
have occurred more commonly on the tree representing the sample of species.

When we controlled for the observed sampling deviations from the random
sampling expectation, our PGLS test of diversification effects on body size did not find
any association between these variables ($\beta = 0.00$, $p = 0.71$). When we removed the
sampling variable from the analysis, however, the PGLS obtained a trend more consistent
with the results of the BiSSE analysis ($\beta = 0.08$, $p = 0.081$). Although this PGLS test
cannot technically distinguish between body-mass related speciation and extinction
effects, we would expect to find a positive association between diversification and body
mass if speciation rate correlates positively with body mass and extinction rates do not
(as suggested by the above tests). Additionally, simulations by Freckleton et al. (2008)
showed that their test was much more sensitive to speciation effects than to extinction
effects. Thus, given the much lower power of the test to detect an effect of extinction,
any positive result would more likely result from speciation effects.

4. DISCUSSION

We tested a series of predictions involving the links among body mass, phylogenetic
diversification, and current extinction risk in primates. With regard to current patterns of
extinction, we found that larger bodied species experience higher extinction risk. This
result replicates previous findings, but for the first time allows for phylogenetic
uncertainty and uses the scaling parameter $\lambda$ to control for the non-Brownian distribution
of variation in IUCN threat status categories (Cardillo et al. 2005; Clauset & Erwin 2008;
Davies et al. 2008). We also tested, for the first time, whether the ordinal nature of
IUCN threat status categories impacts the statistical performance of independent contrasts and PGLS analyses that use it as a dependent variable. Our 1000 phylogenetic simulations show the ordinal coding itself does not produce elevated type 1 error in our primate dataset.

Despite finding a strong association between threat categories and body mass in primates, we failed to find evidence for an historical association between body mass and extinction rates. Instead, in a limited number of tests, we found that speciation rates may increase with body mass, which was opposite to our predictions. These analyses point to an association between larger body mass and higher speciation rate, which contrasts with most predictions, including ours, that speciation rates covary negatively with body mass (see also Liow et al. 2008). After controlling for biased sampling of species in a PGLS model, however, we found no association between diversification rate—a function of speciation and extinction—and body mass.

Previous studies found that intrinsic biological variables can explain a substantial proportion of variation in risk status (Purvis et al. 2000; Cardillo et al. 2005). These previous statistical models found that high risk variables, including large body mass, high trophic level, small geographic range, and slow life history, explain threat status to a substantial degree and independently of anthropogenic effects (Purvis et al. 2000; Cardillo et al. 2005; Purvis 2008). Thus, present day extinction risk status should be related to the historically experienced extinction rates of lineages.

We found it surprising that current patterns of extinction risk covaried with body mass in our study and previous studies, while estimates of extinction rates across primate phylogeny failed to show this effect. It could be that higher threat categories among
large-bodied primates have resulted only from anthropogenic effects in the present.

Alternatively, the tests may have failed to detect differences because available methods to estimate historical extinction rates suffer from low statistical power (Maddison et al. 2007; see also Nee et al. 1992; Kubo & Iwasa 1995; Purvis et al. 1995; Gardezi & da Silva; 1999; Paradis 2004; Rabosky 2010). Using simulations, for example, Maddison et al. (2007) showed that the power to detect variation in speciation rates is relatively high (power for speciation in simulations of 500 species was ~58%), as compared to the power of detecting variation in extinction rates (~21%). Rabosky (2010) recently showed that, in addition to low power, current methods cannot accurately estimate extinction rates when these rates vary across the tree. His analyses were conducted with complete sampling and phylogenetic information of every extant species. Given the fossil evidence for a plethora of extinct primate species, we agree with Rabosky (2010) that future analyses should incorporate fossil information as a means to increase power and to estimate extinction rates more accurately (see also Purvis 2008).

Our study also offers a cautionary tale about the importance of testing the assumptions of methods for studying speciation and extinction rates. When using a new method to account for incomplete sampling of taxa, we found some support for an association between speciation and body mass. However, our data violated another assumption of these methods, namely that sampling of species is random with respect to body mass. When we controlled for non-random sampling, the association between diversification rates and body mass became non-significant. Thus, we offer a valuable empirical example of shortcomings to these methods that complement previous simulation studies, and we provide a procedure for investigating biased sampling for
others to use in future studies. The potential effects of nonrandom sampling are a serious concern for previous studies of trait dependent diversification (Purvis et al. 1995; Paradis 2005; Freckleton et al. 2008; Nakagawa & Freckleton 2008).

We began this study with the prediction that populations of smaller bodied animals may speciate more quickly, as they experience vicariance events more frequently or they are able to better partition the environment into more niches (Gardezi & da Silva 1999). Previous studies provided hints that speciation rate could increase with body mass in primates, rather than showing a strict negative correlation with body mass. For example, Purvis et al. (1995) found that the family Cercopithecidae (Old World monkeys) experienced larger speciation rates than did other primate lineages. While their study did not assess how speciation covaried with characteristics such as body mass, our results are consistent with their findings given that the cercopithecids are generally larger bodied than extant platyrrhine and strepsirrhine primates that more frequently fall below Kay’s Threshold. The cercopithecids also are typically smaller than the (relatively) species depauperate hominoids. These clade-specific patterns could be tested more rigorously with the new statistical software MEDUSA (Alfaro et al. 2009). Our results are consistent with Freckleton et al.’s (2008) findings of a significant correlation between body mass and diversification rate. The results imply that some degree of species selection on body mass may have occurred in primates, in that the influence of body mass on cladogenesis explains the distribution of body mass even after allowing for anagenic change as it is incorporated into the model (Supplemental Figure 1; Vrba 1984; Purvis et al. 1995; Maddison et al. 2007).
In summary, we used new methods to incorporate phylogenetic uncertainty and to assess how body mass influences primate diversification. Our results add to a growing body of evidence that larger bodied animals are more susceptible to extinction. The lack of significance for historical patterns of extinction may indicate that methods to detect extinction rates from extant species are severely compromised in statistical power (Kubo & Iwasa 1995; Purvis et al. 1995; Gardezi & da Silva 1999; Paradis 2004; Maddison et al. 2007; Rabosky 2010). Intriguingly, we found some evidence that speciation rates appear to increase with body mass in primates (see also Liow 2008). It will be interesting to see if similar patterns occur in other groups of vertebrates and, if so, how and why the effects of body size on speciation vary among clades.
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Figure 1. Sliding break point analysis of body mass dependent speciation and extinction rates on the consensus tree. Solid lines connecting points show rates for species that have larger body mass than the given point on the body mass X-axis (black 95% credible interval bars), while dashed lines are for species with smaller body mass (grey 95% credible interval bars). Triangles in Fig. 1a show inferred extinction rates and squares in Fig. 1b show inferred speciation rates. Thick vertical arrows correspond to our three \textit{a priori} hypotheses.
Table 1. Fossil calibration ranges used to date molecular phylogenies

<table>
<thead>
<tr>
<th>MRCA node</th>
<th>Min. Age (ma)</th>
<th>Max. Age (ma)</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Homo-Pan</em></td>
<td>5</td>
<td>8</td>
<td>Haile-Selassie (2001)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Senut <em>et al.</em> (2001)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Vignaud <em>et al.</em> (2002)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Brunet <em>et al.</em> (2002)</td>
</tr>
<tr>
<td><em>Homo-Pongo</em></td>
<td>12.5</td>
<td>18</td>
<td>Kelley (2002)</td>
</tr>
<tr>
<td><em>Papio-Theropithecus</em></td>
<td>3.5</td>
<td>6.5</td>
<td>Leakey (1993)</td>
</tr>
<tr>
<td>extant Catarrhini</td>
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<td>30.0</td>
<td>Young &amp; MacLatchy (2004)</td>
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<td></td>
<td></td>
<td></td>
<td>Benefit &amp; McCrossin (2002)</td>
</tr>
<tr>
<td><em>Cebus-Saimiri</em></td>
<td>12.5</td>
<td>NA</td>
<td>Hartwig &amp; Meldrum (2002)</td>
</tr>
</tbody>
</table>
Supp. Fig. 1. Sliding break point analysis of body mass dependent transition rates on the consensus tree. Solid lines connecting points show rates for species that have larger body mass than the given point on the body mass X-axis (black 95% credible interval bars), while dashed lines are for species with smaller body mass (grey 95% credible interval bars). Thick vertical arrows correspond to our three a priori hypotheses.