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Modeling lineage and phenotypic diversification in the New World monkey (Platyrrhini, Primates) radiation

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1	Modeling	lineage and	phenotypic	diversification	in the Nev	w World monkey

- 2 (Platyrrhini, Primates) radiation
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- 17 Short title: Modeling platyrrhine diversification

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1 ABSTRACT

2 Adaptive radiations that have taken place in the distant past can now be more 3 thoroughly studied with the availability of large molecular phylogenies and comparative 4 data drawn from extant and fossil species. Platyrrhines are a good example of a major 5 mammalian evolutionary radiation confined to a single continent, involving a relatively 6 large temporal scale and documented by a relatively small but informative fossil record. 7 Here, we present comparative evidence using data on extant and fossil species to 8 explore alternative evolutionary models in an effort to better understand the process of 9 platyrrhine lineage and phenotypic diversification. Specifically, we compare the 10 likelihood of null models of lineage and phenotypic diversification versus various models of adaptive evolution. Moreover, we statistically explore the main ecological 11 12 dimension behind the platyrrhine diversification. Contrary to the previous proposals, 13 our study did not find evidence of a rapid lineage accumulation in the phylogenetic tree 14 of extant platyrrhine species. However, the fossil-based diversity curve seems to show a 15 slowdown in diversification rates toward present times. This also suggests an early high 16 rate of extinction among lineages within crown Platyrrhini. Finally, our analyses support 17 the hypothesis that the platyrrhine phenotypic diversification appears to be characterized 18 by an early and profound differentiation in body size related to a multidimensional 19 niche model, followed by little subsequent change (*i.e.*, stasis). 20 *Keywords*: body size; adaptive radiation; fossil record; niche-filling 21

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1 1. Introduction

2 The study of lineage and phenotypic diversification in living clades with 3 relatively recent divergences has supported models of adaptive radiation (Simpson, 4 1953; Schluter, 2000) that predict selective influences arising from ecological 5 opportunity and circumstances (Gavrilets and Losos, 2009; Losos and Mahler, 2010) 6 This process frequently results in great phenotypic variation and species richness 7 relative to a short time frame of phylogenetic divergence (Schluter, 2000; Gavrilets and 8 Losos, 2009; Losos and Mahler, 2010). It is also suspected that similar processes of 9 adaptive radiation have driven the initial diversification of clades that originated in the 10 distant past, for example, at the ordinal and subordinal levels in mammalian clades 11 (Gavrilets and Losos, 2009; Losos and Mahler, 2010). In such cases, ecological 12 opportunity may decrease during the radiation as niches are filled, leading to stasis. 13 Several evolutionary radiations have been carefully studied with recently 14 developed mathematical models applied to molecular phylogenies and comparative data 15 from extant species (Nee, 2006; Gavrilets and Losos, 2009; Losos and Mahler, 2010). 16 However, it is becoming clear that our capacity to successfully model and understand 17 the diversification processes of ancient clades is limited when only neontological data 18 are considered (Quental and Marshal, 2010; Slater et al., 2012). New World monkeys 19 (Parvorder Platyrrhini), one of the three major groups of living and fossil primates, are a 20 good example of a major mammalian evolutionary radiation that occupies a large 21 temporal scale (*i.e.*, 20-40 million years ago or megannums [Ma] in Central and South 22 America), exhibits a remarkable phenotypic variation (e.g., a body mass spanning two 23 orders of magnitude, from 0.1 to more than 10 kg) and presents a relatively small but 24 informative fossil record (Fleagle, 1999; Fleagle & Tejedor, 2002; Tejedor, 2008).

1	Morphological and phylogenetic studies have hypothesized that the diversification of
2	this monophyletic group was mainly linked to the action of deterministic-selective
3	factors related to ecological variables (Rosenberger, 1992; Marroig and Cheverud, 2001;
4	Rosenberger et al., 2009). However, there is no general agreement about the main
5	ecological dimension — e.g., diet, locomotion or a multidimensional niche— behind the
6	platyrrhine diversification (Rosenberger, 1992; Allen and Kay, 2011; Youlatos and
7	Meldrum, 2011; Perez et al., 2011). It has also been suggested that the marked
8	phenotypic diversification of platyrrhines occurred relatively quickly during the initial
9	branching process of the main extant clades in connection with ecological niche
10	opportunity (i.e., an early-burst platyrrhine radiation), followed by a slowdown in
11	evolutionary rates which resulted in the widespread retention of the formative patterns
12	that are characteristic of these lineages (<i>i.e.</i> , evolutionary stasis; Rosenberger, 1992;
13	Rosenberger et al., 2009; Perez et al., 2011).
14	Although some recent studies of evolutionary radiations have included
15	information from the fossil record along with phylogenies and comparative data of
16	extant species in a mathematical modeling framework (Slater et al., 2010, 2012; Etienne
17	et al., 2011), such approach to the study of the dynamic processes of lineage origin and
18	extinction, and phenotypic diversification, has not been applied to explore the
19	platyrrhine evolutionary radiation. Here, we present comparative evidence using data on
20	extant and fossil species to explore alternative evolutionary models in an effort to better
21	understand the process of platyrrhine lineage and phenotypic diversification.
22	Specifically, we compare the likelihood of null models of lineages and phenotypic
23	diversification versus various models of adaptive evolution. Moreover, we explore the
24	main ecological dimension behind the platyrrhine diversification. If the platyrrhine

1 diversification conforms to the adaptive radiation theory, we expect that differentiation 2 of the major extant platyrrhine lineages was concentrated relatively early in the history 3 of the clade and that phenotypic variation — measured as body size— was partitioned 4 among subclades early in their phylogenetic history as well, as a major driver or 5 consequence of ecological niche partitioning and niche-filling. As a starting point, we 6 first estimate a chronophylogenetic tree for most extant platyrrhine species using 7 molecular data and Bayesian methods (Drummond et al., 2006). Then, using this tree 8 and comparative statistical methods we explore the pattern of lineage diversification 9 through time (Nee et al., 1992; Harmon et al., 2003; Ricklefs, 2007; Stadler, 2011a), investigate the pattern of body size diversification through time and the fit of a series of 10 11 evolutionary models (Harmon et al., 2003; Butler and King, 2004). Finally, given that 12 the inference of the tempo and mode of diversification of a clade using only extant 13 species can be biased (Quental and Marshal, 2010; Slater et al., 2012), we compare the 14 results based on extant species with the estimated body masses and number of lineages 15 inferred from the platyrrhine fossil record. Summarizing, our work contributes to the 16 discussion of platyrrhine evolution and diversification in three different ways: 1) we 17 present one of the most complete molecular phylogenies of extant platyrrhine species to 18 date, sampling published data on 108 taxa and estimating a chronophylogenetic tree for 19 78 putative "good" species; 2) we mathematically model the pattern of lineage and 20 phenotypic diversification of the platyrrhine clade; and 3) we combine data about extant 21 and fossil species in a novel way not employed before in studies of the platyrrhines to 22 better understand the process of their diversification.

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1 **2. Materials and methods**

2 2.1. Molecular divergence among species and phylogenetic inference

3 Phylogenetic trees are used in almost every branch of evolutionary biology. In 4 particular, in comparative analyses they are needed to avoid misinterpreting historical 5 contingencies as causal relationships and to understand the patterns of diversification 6 (Nee, 2006; Losos, 2011; Yang and Rannala, 2012). In our study, the phylogenetic tree 7 itself is not of direct interest but it is a necessary first step for the following statistical 8 comparative analyses. Previous phylogenetic studies have estimated the platyrrhine 9 phylogenetic tree using ca. 50% (58 species, Chatterjee et al., 2009; or 64 species, 10 Perelman et al., 2011) or 60% (73 species, Fabre et al., 2009) of species recognized by 11 Groves (2005). However, most of the methods in comparative analyses assume that a 12 large fraction of the extant species in the clade under study are included in the dataset (Pybus and Harvey, 2000; Ricklefs, 2007; Cusimano et al., 2012). On the other hand, 13 14 the use of many different definitions of species in primate systematics has led to a 15 fluctuating taxonomy in platyrrhines (e.g. Groves, 2004, Rylands et al., 2011) and 16 concerns on taxonomic inflation have arisen (Rosenberger, 2012). Since speciation 17 events are not instantaneous, the decision on to how finely distinguish lineages is rather 18 arbitrary (Ricklefs, 2007). For this reason, we decided to set up a reproducible criterion 19 for distinguishing species based on a minimum molecular divergence threshold. A total 20 of 108 platyrrhine species and subspecies were initially considered for analysis based on 21 the availability of DNA sequences on the Genbank database (Supplementary Table A.1). 22 Cytochrome b (CytB) DNA sequences were downloaded from Genbank for 90 23 specimens (based on availability) that were previously accepted as valid species or 24 subspecies in the platyrrhine literature (Groves, 2005). We estimated molecular distance

1 among the 90 specimens within each genus using Mega 5 (Tamura et al., 2011) and 2 established a likely point (3%) of minimum divergence among species (Fouquet et al., 3 2007; Clemente-Carvalho et al., 2010). After this procedure, we excluded taxa with low 4 molecular divergence (mainly subspecies) and considered as full species those taxa 5 previously regarded as subspecies that were above the distance threshold. Specifically, 6 26 taxa were excluded from analysis based on the existence of a low molecular distance, 7 because they probably represent geographic variants of other species included in the 8 dataset (Supplementary Table A.1). The final dataset included 78 species, as 14 taxa 9 from the Perelman et al. (2011) dataset that did not have an available CytB sequence 10 were included based on the fact that they are broadly recognized as full species. Another 11 four species were excluded because their large proportion of missing data could hinder 12 the analysis. Thus, our dataset comprised a great percentage (ca., 80%) of the likely 13 extant platyrrhine species.

14 To estimate the chronophylogenetic tree for the 78 platyrrhine species we 15 analyzed the dataset obtained for the platyrrhine species and three outgroups (Macaca 16 mulatta, Pan troglodytes, Homo sapiens) from GenBank (see Wildman et al., 2009, and 17 Perelman et al., 2011). The analyzed dataset is a DNA concatenated matrix for 81 18 species with a total of 25,361 bp, including 15 selected nuclear genes from Perelman et 19 al. (2011) with few missing data, and 11 non-coding sequences from Wildman et al. 20 (2009) (Supplementary Table A.2). The dataset also includes IRBP and ξ -globin genes 21 and four mitochondrial sequences (CytB, 16s, 12s and COxII). Several available gene 22 sequences (e.g. from Perelman et al., 2011) were not included in our phylogenetic 23 estimation because a great number of species lacked data for those genes. The 24 sequences of each gene were aligned using ClustalW and manually corrected with

BioEdit 7.0.0 software (Hall, 2004). Accession numbers for the mitochondrial, IRBP
 and ξ-globin sequences are shown in Supplementary Table A.1.

3 We performed a concatenated Bayesian analysis based on the molecular dataset 4 to estimate chronophylogenetic relationships among the 81 primate species. jModelTest 5 0.1 (Posada, 2008) was employed to determine the most appropriate model of sequence 6 evolution for each analyzed gene sequences estimated under the Akaike Information 7 Criterion with correction for sample size (AICc). The best fit models for studied genes 8 are shown in the Supplementary Table A.2. The Bayesian chronophylogenetic analysis 9 was performed using BEAST v1.6.1 (Drummond and Rambaut, 2007). The analysis was 10 carried out using Markov Chain Monte Carlo (MCMC) simulations for 200,000,000 11 generations and a sample frequency of 20,000. The convergence was determined using 12 the program Tracer v1.5 (Rambaut and Drummond, 2007) and the first 1,250 sampled 13 trees were excluded. We used a relaxed molecular clock model, which allows 14 substitution rates to vary across branches according to an uncorrelated lognormal 15 distribution (Drummond et al., 2006; Drummond and Rambaut, 2007). Six fossil 16 calibrations were selected (see Perez et al., 2013). Fossil calibration for Homo-Pan 17 divergence (minimum time 5.7 Ma, maximum time 10 Ma; LogNormal distribution 18 with offset 5.7, mean 0.5 and standard deviation 0.5) and Anthropoidea (minimum time 19 33.70 Ma, maximum time 65.80 Ma; LogNormal distribution with offset 33.7, mean 2.4 20 and standard deviation 0.55) were obtained from Benton et al. (2009). Minimum 21 divergence time of Alouattinae, Cebinae and Aotinae were set at 12.5 Ma, based on 22 Stirtonia, Neosaimiri and Aotus dindensis, three fossils attributable to Alouattinae, 23 Cebinae and Aotinae, respectively (Kay et al., 2008; Tejedor, 2008). We used a 24 LogNormal distribution with offset 12.5, mean 1.8 and standard deviation 0.4 for the

1	Alouattinae, Cebinae and Aotinae constraints. Minimum divergence time of Pitheciidae
2	was set at 15.7 Ma (LogNormal distribution with offset 15.7, mean 1.5 and standard
3	deviation 0.5), based on Proteropithecia neuquenensis, a fossil attributable to
4	Pitheciidae (Kay et al., 1998). Maximum divergence time was set at 26 Ma, based on
5	the absence of Pitheciidae, Alouattinae, Cebinae and Aotinae fossils in or previous to
6	the Deseadan fauna of Salla, Bolivia, and other South American formations of the same
7	age. Both minimum and maximum calibration bounds were set to the probability that
8	the true divergence time outside the bounds is small, but non-zero (dos Reis et al., 2012;
9	Perez et al., 2013). Although a calibrated tree could be important for discussing some
10	aspects of the results, the following analyses are insensitive to the total length of the
11	tree, <i>i.e.</i> based on relative times (see discussion below). We computed the maximum
12	clade credibility (MCC) tree in TreeAnnotator 1.4.8 (Drummond and Rambaut, 2007).
13	FigTree v1.3.1 was used to plot the phylogenetic tree.

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15 2.2. Lineage diversification pattern analysis

16 The number of species in a clade is the result of speciation and extinction 17 processes acting since the origin of that clade. The most direct way of studying this 18 diversity dynamic is to analyze the fossil record. However, the information contained in 19 molecular phylogenetic trees of extant species allow us to estimate speciation and 20 extinction rates in the absence of an appropriate fossil record (Harvey et al, 1994; Nee et 21 al, 1994; Ricklefs 2007; Stadler, 2011a). This information can be recovered by 22 analyzing the fit of different mathematical models of diversification to the reconstructed phylogeny. 23



Following this approach, we first quantitatively tested, using the widely used γ

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1	statistic (Pybus and Harvey, 2000), whether the pattern of lineage diversification in the
2	platyrrhine tree departs from what is expected under the most simple model, Pure-Birth
3	(PB), in which each species has a constant probability λ (speciation rate) of generating
4	another species in each point in time and there is no extinction (Nee, 2006). Under this
5	model, the number of species N(t) after t units of time, starting from N(o) species is
6	expected to grow exponentially: $N(t) = N(o)e^{\lambda t}$. The γ statistic measures the difference
7	between the average sum of branch lengths between each internal node and the root and
8	the midpoint of the tree (Pybus and Harvey, 2000). This statistic has mean of 0 for trees
9	generated under a PB process, and significant negative values indicate a decelerating
10	lineage accumulation rate toward present times, or, in other words, that branching
11	events are concentrated disproportionally early in the phylogeny, as is expected in
12	scenarios of adaptive radiation (Gavrilets and Losos, 2009).
13	To further assess the diversification model that best explains the observed
14	diversity pattern and to estimate possible changes in diversification rates across the
15	platyrrhine tree we compared the fit, using the Akaike Information Criterion (AIC), of
16	the observed branching sequence to various models of lineage accumulation. AIC is
17	calculated as $-2\ell+2K$ where ℓ is the maximum likelihood value of the data and K is the
18	number of parameters in the model. AIC represents a compromise between fit and
19	complexity of the model (Burnham and Anderson, 2002). The tested models were the
20	following: (i) PB and birth-death (BD; with parameters λ and μ [extinction rate]), as
21	constant-rates lineage accumulation models (although some of these models allow for
22	discrete λ shifts at specific points in time), and (ii) linear (DDL) and exponential (DDX)

density-dependent processes (as variable λ models with no extinction) (Rabosky and

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24 Lovette, 2008a). Because the importance of extinction in shaping lineage origination

1	curves may be underestimated in the DDX and DDL models, we also included a
2	density-dependent model with a non-zero μ (DDD+E; Etienne et al., 2011). Density-
3	dependent models assume that the diversification rate $(\lambda - \mu)$ decreases as the lineage
4	population reaches some threshold density. If there are ecological limits to diversity, a
5	slowdown in lineage diversification rate would be expected in an adaptive radiation
6	scenario, since ecological opportunity may decrease as niches are filled (Rabosky and
7	Lovette, 2008b; Gavrilets and Losos, 2009; Ettiene et al., 2011). Also, since λ and μ
8	may have changed through time in response to different external factors, we also
9	included the recently proposed birth-death-shift (BDS) method by Stadler (2011b),
10	which can estimate whether and when a shift in rates occurred in the tree.
11	To account for incomplete sampling in our tree, we added the missing splitting
12	events to the analyzed tree, using the maximum likelihood λ and μ estimations from the
13	MCC tree, assuming that they occurred between 1 and 5 Ma. Using this approach, we
14	simulated 200 completely sampled trees and repeated the γ test and the model fitting
15	analyses (Cusimano et al., 2012). To also account for uncertainty in phylogenetic
16	reconstruction, analyses were repeated over a random sample of 100 trees drawn from
17	the post-burn in set of the BEAST analysis.
18	Finally, we visually inspected the time pattern of lineage accumulation by
19	plotting the number of ancestral lineages in our phylogenetic tree (in a log scale) versus
20	time (lineage-through-time plot, LTT; Ricklefs 2007), and compared it, using
21	simulations, to the expected pattern under the PB and BD (keeping track of extinct
22	lineages) models of diversification.
23	All analyses were conducted with the LASER (Rabosky, 2006), TreePar and
24	TreeSim (Stadler, 2011b,2011c), and APE (Paradis et al. 2004) packages for R 2.15.1

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1 (R-Development Core Team, 2012).

2	In order to develop a generic origination trajectory for the fossil record, we
3	assembled geochronologic ages from the literature for 23 fossil platyrrhine genera
4	ranging from the Deseadan South American Land Mammal Age (SALMA) to the
5	Laventan SALMA, spanning from 26 to 11.8 Ma (Fleagle and Tejedor, 2002; Tejedor,
6	2008; Rosenberger et al., 2009). This data was then used to build origination estimates
7	(as numbers of genera confined to an interval plus those that cross the top boundary of
8	the interval; Foote, 2000) for each SALMA. To test for possible biases due to a poor
9	sampling of the fossil record, we applied the Spearman's rank correlation test as
10	described in Barrett et al. (2009). Briefly, the number of genera for each time slice (as
11	SALMAs) is expected to be positively correlated with the number of sampled
12	geological formations if sampling is affecting the shape of the origination curve.
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24 and female data were pooled and log-transformed for all analyses.

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1	To explore the time pattern of body mass variation, we first calculated a mean
2	relative disparity-through-time (DTT) plot for our phylogeny as described by Harmon et
3	al. (2003). Disparity is measured as $D = \Sigma(d_i)/n-1$ where d_i is pairwise Euclidean
4	distance between species and n is the number of species. First, disparity was calculated
5	for the entire platyrrhine clade, and then for each subclade. Disparity of each subclade
6	was standardized by dividing it by the disparity of the entire clade (relative disparity;
7	Harmon et al., 2003). Finally, the mean relative disparity for each point in time is
8	calculated for all subclades present at that time. Then we compared the observed body
9	mass disparity trough time to that expected if character evolution had followed a
10	Brownian motion (random) model of diversification by simulating body size evolution
11	1,000 times across our tree. Disparity values near 0 imply that most of the phenotypic
12	variation is partitioned among subclades rather than within each subclade (e.g. among
13	families, subfamilies, etc. rather than within each family, subfamily, etc.), whereas
14	values near 1 imply the opposite, indicating that subclades have independently evolved
15	to occupy similar places of morphological space (Harmon et al., 2003). Phenotypic
16	disparity is expected to be partitioned among subclades early in an early niche-filling
17	scenario of adaptive radiation. We also calculated the morphological disparity index
18	(MDI, Harmon et al., 2003), which quantifies the overall difference between the
19	observed and expected curves of disparity through time. Negative values are expected
20	under an adaptive radiation scenario, indicating lower subclade disparity than in the
21	random evolution model. The DTT analysis is implemented in the GEIGER package
22	(Harmon et al., 2008) for R.

Body mass is proposed to be strongly related to diet among platyrrhines
(Rosenberger, 1980, 1992; Marroig and Cheverud, 2001; Perez et al., 2011). To explore

1 this hypothesis, we applied the model selection approach of Butler and King (2004) to 2 compare the relative fit (measured with Akaike Information Criterion, AIC; see above) 3 of a stochastic model (Brownian motion, BM; Felsenstein, 1985) vs. seven models of 4 adaptive evolution (Ornstein-Uhlenbeck, OU; Hansen, 1997) to the body size variation. 5 In the BM model, body size evolves up the phylogeny via random walk and disparity 6 accumulates over time (Felsenstein, 1985). To model adaptive evolution in platyrrhine size variation, we implemented seven Ornstein-Uhlenbeck models (OU; Hansen, 1997) 7 8 with either one, three, four or five optima. Particularly, we implemented a random walk 9 with a single stationary peak modeled as an Ornstein–Uhlenbeck process (OU1), such 10 that the size have a tendency to return to a median value (Hansen, 1997; Butler and 11 King, 2004), and six OU models with several optima formulated based on previous 12 hypotheses about the main ecological dimension behind the platyrrhine diversification. The parameters of the evolutionary models were estimated by maximum likelihood: σ 13 14 (the intensity of the random changes in body size), α (the rate of changes toward an 15 optimum, or the strength of selection), θk (optimal value for the body size in each niche 16 optimum k). 17 To build the ecological models, or adaptive evolution hypotheses, we first 18 assigned each extant species to an ecological niche according to published works 19 (Rosenberger, 1992; Norconk et al., 2009; Youlatos and Meldrum, 2011; Allen and Kay,

20 2011), which describe different ecological groups. For uni-dimensional niche
21 hypotheses, data concerning diet composition (i.e., average annual percentages of plant
22 parts and insects in the diets of platyrrhine genera), diet quality (i.e., percentage of
23 structural plant parts [leaves and stems], reproductive plant parts [fruits, seeds, flowers,
24 nectar and gums] and animal matter in the diet) and locomotion (i.e., percentages of

1 arboreal quadrupedal walk, clamber and bridge, clawed locomotion and suspensory 2 locomotion) of platyrrhine genera and species were taken from Norconk et al. (2009), 3 Allen and Kay (2011) and Youlatos and Meldrum (2011), respectively. We then 4 calculated the principal component (PC) scores for each data set to reduce the number 5 of ecological variables and avoid multicollinearity; these PCs describe broad variation 6 in ecology (i.e., diet composition, diet quality and locomotion PCs) and were used to 7 group species in the different ecological niches (see below). It is important to remark 8 that although these niche models are based on real datasets, they are hypotheses 9 modelling the ancestral partition of platyrrhine niches. OU processes model the effect of 10 different selective regimes (ecological niches in this case) acting along the branches of a 11 phylogenetic tree and thus can be used to test for phenotypic diversification related to 12 ecological factors (Buttler and King, 2004), using data that is strongly phylogenetically 13 structured. In our analyses, character states for all internal branches were estimated 14 using a maximum likelihood approach. Finally, we explored body mass variation for the 15 ancient Patagonian and younger La Ventan (Colombia) fossil species in relation with the 16 best supported model. The OU analysis is implemented in the OUCH package (Buttler 17 and King, 2004) for R and was performed using the function OU average from Jaffe et 18 al. (2011).

To account for uncertainty in phylogenetic reconstruction, DTT and OU analyses
were performed over a random sample of 100 trees drawn from the post-burn in set of
the BEAST analysis.

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23 **3. Results**

Our chronophylogenetic tree of platyrrhines is in general agreement with other

1	recent relationship estimations (Fig. 1; Opazo et al., 2006; Wildman et al., 2009;
2	Perelman et al., 2011; Perez et al., 2012), which supports the division of platyrrhines
3	into three monophyletic families (Atelidae, Cebidae and Pitheciidae) and suggests a
4	sister-group phylogenetic relationship between Atelidae and Cebidae (Opazo et al.,
5	2006; Wildman et al., 2009). We found <i>Aotus</i> to be phylogenetically related to
6	Callitrichinae, as did Perelman et al. (2011), but with low node support. The affinities of
7	Aotus are contentious as several molecular and morphological studies do not align
8	(Perez et al., 2012; Rosenberger and Tejedor, 2013). Because of this controversy, we
9	repeated all the statistical analyses varying the phylogenetic position of Aotus (as a
10	sister clade of Cebidae or Cebinae, as most studies report). The results were almost
11	identical (data not shown). Our results are also consistent with other molecular
12	assessments, which indicate the branches connecting the main platyrrhine lineages are
13	short, suggesting that the diversification of the main living platyrrhine clades could
14	indeed represent a rapid sequential radiation (but see below; Fig. 1; Opazo et al., 2006;
15	Wildman et al., 2009; Perez et al., 2012). The divergence time estimation suggest that
16	the last common ancestor (LCA) of extant platyrrhine primates existed at ca. 25 Ma,
17	with the 95% confidence limit for the node ranging from <i>ca</i> . 22.5–29 Ma. This result is
18	in contrast with other recent estimations, mainly to the influential Hodgson et al. (2009)
19	study, which suggested a LCA for crown platyrrhines at ca. 19 Ma.
20	Based on the γ statistic over 100 random trees drawn from the post-burn in set,
21	we found no evidence of a decelerating lineage accumulation rate toward present times
22	in the overall origination pattern of the platyrrhine clade (mean $\gamma = 0.493$, $p = 0.679$).
23	This result is robust to incomplete sampling (mean $\gamma = 0.290$, $p = 0.612$). Moreover, the
24	best supported model of lineage diversification was a pure-birth model with one rate

1 shift: an abrupt slowdown in speciation rates at 0.42 Ma (Table 1). Such recent rate 2 shifts may not be significant since incomplete species sampling and taxonomic inflation 3 can affect the shape of the most recent portion of the tree; however, when accounting for 4 incomplete sampling, a slowdown is also recovered at similar dates (Supplementary 5 Table A.4). Overall, this result shows, based on the extant species tree, that through 6 more than 99% of the time since its origins, crown platyrrhines diversified at a constant 7 rate. On the other hand, for the BD models, the maximum likelihood extinction rate is 8 probably underestimated, an acknowledged issue of diversification models. The absence 9 of an evident early pulse of lineage origination is also visually seen in the LTT plot for 10 extant species (Fig. 2). However, the fossil lineages origination curve seems to show a 11 different picture: a relatively rapid increase in the number of species followed by an 12 apparent slowdown in the origination rate, a pattern reminiscent of a density-dependent 13 diversification trajectory (Fig. 2, blue dashed line). This pattern is even stronger when 14 compared to simulated trees under the PB and BD models, even considering that the BD 15 curve includes extinct lineages (Fig. 2). Furthermore, this minimum estimated diversity 16 through time is likely to be very conservative, since it lacks a large fossil record from 17 the early Miocene of Amazonia, an area that was certainly populated with platyrrhine 18 representatives. The Spearman's correlation coefficient between genera per SALMAs 19 and the sampled geological formations showed a non-significant value (r=0.286, p=20 0.65), suggesting that the observed increase in fossil genera over time is not due to 21 sampling bias.

Figure 3 shows the DTT plot for the body mass data. Average subclade disparity along the entire history of the group is lower than expected under a BM model of mass evolution. Values drop near 0 since the early divergence of the platyrrhines, and show

1	little variation over time. An MDI value of -0.255 also confirms quantitatively this
2	result. This outcome indicates a strong pattern where most size variation occurs among
3	the main platyrrhine subclades, which tend to occupy more isolated regions of body
4	mass morphospace, consistent with an early radiation and niche-filling scenario.
5	The PCs scores displayed in the figure 4 were used to group species in the
6	different ecological niches. Particularly, we defined 4 and 3 niches for diet composition
7	(OU-dietC4 and OU-dietC3, respectively); 5 and 4 niches for diet quality (OU-dietQ5
8	and OU-dietQ4, respectively); and 3 niches for locomotion (OU-Loc3; Fig. 4). Finally, a
9	multidimensional niche model was built based on the Rosenberger (1992) hypothesis;
10	this hypothesis is mainly a combination of diet composition and locomotion niche
11	dimensions and defines 5 broad ancestral ecological niches (OU-MD5; Fig. 4). The
12	overall fit of the models of body mass evolution to these hypothetical niches is shown in
13	Table 2. The OU model with 5 body mass optima (OU-MD5), following the
14	multivariate niche hypothesis of Rosenberger (1980, 1992), was the best supported, with
15	an Akaike weight well above the other candidate models, which performed poorly in
16	comparison. Taking into consideration the proposed phylogenetic relationships of the
17	fossils with extant clades (Rosenberger, 1992; Fleagle, 1999; Fleagle and Tejedor, 2002;
18	Rosenberger et al., 2009), body mass estimates for fossil platyrrhines, particularly La
19	Venta genera, show a similar pattern of variation to that observed for each of the
20	ecological categories of the OU-MD5 model (Fig. 5).
21	

22 **4. Discussion**

The process of diversification of the platyrrhines has been investigated
phylogenetically and morphologically (*e.g.*, Rosenberger, 1992; Fleagle, 1999; Marroig

1	and Cheverud, 2001; Hodgson et al., 2009; Wildman et al., 2009; Perez et al., 2011,
2	2012, 2013). Many of these studies have interpreted the diversification of extant New
3	World monkeys as an adaptive radiation in which the major lineages diversified early
4	and into various alternative ecological niches that continue to exist today (see
5	discussion in Rosenberger 1992; Rosenberger et al. 2009; Tejedor, 2012). In all cases,
6	the initial time of the diversification process of extant platyrrhine species is emphasized
7	(e.g., Kay et al., 2008; Hodgson et al., 2009; Rosenberger, 2012; Perez et al., 2013).
8	Previous works suggest that diversification of platyrrhines conforms to key expectations
9	of a model of adaptive radiation —diversity or time-dependent lineage origination
10	associated with phenotypic diversification of ecologically relevant traits such as body
11	size. However, to our knowledge, this is the first statistical effort to model the
12	phenotypic and lineage diversification of platyrrhines using a large and representative
13	species sample and combining molecular phylogenies, fossil data and a comparative
14	phylogenetic approach. In the following sections, we discuss our results on the pattern
15	of lineage branching and phenotypic diversification, its relationships with previously
16	hypothesized ecological variables and the importance of the previously suggested initial
17	times of branching to understand this diversification process.

18

19 4.1. Lineage diversification

As it has been suggested often for platyrrhines (*e.g.*, Hodgson et al., 2009; Wildman et al., 2009), a central prediction of the adaptive radiation model based on young radiations is that a large number of lineages are accumulated during the early stages of a clade's evolutionary history, followed by a slowdown in species origination rates (Gavrilets and Losos, 2009; Losos and Mahler, 2010). Contrary to the previous

1 proposals, our study did not find evidence of a rapid lineage accumulation in the 2 phylogenetic tree of extant platyrrhine species. However, it has been repeatedly argued 3 that high extinction rates and other factors might erase the signal of a decrease in 4 lineage diversification rates estimated from phylogenies based on extant species, and 5 that including information from the fossil record is thus essential to account for this 6 (Pybus and Harvey, 2000; Rabosky and Lovette, 2008b; Quental and Marshall, 2010; 7 Slater et al., 2012). In this sense, in our study the fossil-based diversity curve seems to 8 show a slowdown in diversification rates toward present times (Fig. 2). When the oldest 9 known fossil lineages (mostly from Patagonia) are considered, the number of lineages 10 increases noticeably compared to those inferred to exist at that time from the molecular 11 phylogeny, and an early pulse (approximately between 20 and 15 Ma) in the origination 12 of platyrrhine lineages may become observable (Fig. 2). Moreover, diversity levels 13 through time estimated from the fossil record are likely underestimated. On the other 14 hand, the absence of a meaningful correlation between the number of fossil genera and 15 platyrrhine-bearing rock formations suggests that this pattern is paleontologically robust 16 and that the platyrrhines present a small but representative fossil record. The observed 17 pattern in the extant diversity curve may be linked to the extinction of the Patagonian 18 primates or their retraction towards the northwest after 15 Ma (Middle Miocene; 19 Tejedor, 2012). Platyrrhines are climate-sensitive mammals that certainly responded to 20 the paleoenvironmental changes in South America after the Middle Miocene, when the 21 environments shifted towards more arid and cooler conditions in Patagonia and the 22 current configuration of Andes and Amazonian originated, generating the extinction of 23 numerous lineages within (Rosenberger et al. 2009) or outside the extant main clades 24 (Kay et al., 2008). Globally, these findings may suggest that high extinction levels

1 played a key role in shaping the extant assemblage of platyrrhine species, and that the 2 observed lineage diversity pattern can be reconciled with the predictions of the adaptive 3 radiation theory only when considering that a significant part of the early platyrrhine 4 diversity became extinct. On the other hand, the observed slowdown in diversification 5 rates near present times, even when incomplete taxa sampling was considered, may be 6 linked to our inability to detect recent lineage splitting events. In this sense, this rate 7 slowdown at Pleistocene times has also been recovered in a recent analysis of a 8 phylogeny of all primates (Springer et al., 2012) in which different taxonomic 9 arrangements were considered, thus further indicating that evolutionary relevant 10 splitting events may be overlooked by our current taxonomic criteria. 11

12 4.2. Phenotypic diversification

13 Our results concerning the temporal pattern of body mass variation are generally 14 consistent with the expectation that the size variation was partitioned among subclades 15 early in the phylogenetic history of the platyrrhines. The plot in the Figure 3 shows how 16 size disparity is high during the early branching process, probably related to changes in 17 ecological conditions, such as ecological opportunity (Harmon et al., 2003). Strikingly, 18 the magnitude of size disparity during the early branching process of platyrrhines is 19 unusually high compared with most previous studies we are aware of (e.g., Harmon et 20 al., 2003; Slater et al., 2010; Derryberry et al., 2011; Weir and Mursleen, 2012), 21 confirming the distinctiveness of the platyrrhine radiation (Delson and Rosenberger, 22 1984).

The AIC analysis used to test whether body mass evolved according to a
 stochastic model or to the occupation of different ecological niches shows interesting

1	results. Particularly, Brownian motion, the non-adaptive and simplest (i.e., with fewer
2	parameters) model, has limited support (Table 2). The models of phenotypic
3	diversification within each dietary and locomotion niche (OU-dietQ, OU-dietC and OU-
4	Loc3 models, Fig. 4) exhibit poor performance with respect to other models (Table 2).
5	The model of phenotypic diversification during differentiation of the main lineages and
6	subsequent evolutionary stasis within the multidimensional ecological niches (OU-MD5
7	model; Rosenberger, 1992; Rosenberger et al., 2009) has the best performance (Table
8	2). These results differ from previous works pointing out that platyrrhine size
9	diversification is mainly related to diet variation (e.g., Marroig and Cheverud, 2001;
10	Perez et al., 2011). Conversely, we show that the diet ecological dimension alone is not
11	enough to explain the platyrrhine body mass diversification. These outcomes support a
12	more complex scenario where platyrrhine evolution is likely related to size changes
13	among the main lineages linked to a multidimensional niche (Rosenberger, 1980, 1992).
14	Moreover, size variation among the platyrrhine fossil species show a similar
15	pattern of variation —particularly the most complete La Venta fossil assemblage— to
16	that obtained analyzing the extant species. This supports the notion of a niche filling at
17	La Venta times, as extant species body size partitioning is already evident in ancient
18	lineages. However, the interpretation of this result is highly dependent on the
19	assignment of fossil species to the extant platyrrhine clades (see below). Taken together,
20	this results may be suggestive of the early platyrrhine lineages diversifying in simpatry,
21	as intraclade competition probably constrained body size evolution.
22	Functional studies pointed out that there are natural size thresholds dividing the
23	platyrrhine dietary-locomotory niches (Rosenberger, 1980, 1992; Fleagle, 1999;
24	Youlatos and Meldrum, 2011). The causal relationship among these variables is debated;

1	for example, Hershkovitz (1977) suggested that the platyrrhine radiation was an
2	evolution of body size, with locomotor and dietary consequences, while other authors
3	suggested the inverse relationship (e.g., Perez et al. 2011). In any case, a central
4	prediction of the adaptive radiation hypothesis is that phenotypes diversify early in the
5	branching process in connection with ecological dimensions (Schluter, 2000; Losos and
6	Mahler, 2010). Thus, the ecological opportunity that could have existed during the early
7	phylogenetic history of platyrrhines within South America probably was a very
8	important factor promoting body size changes among the main lineages; after the initial
9	great diversification, these lineages probably maintained relatively stable size classes
10	linked to their ecological niches. Although ecological opportunity might have driven the
11	early size changes, and narrowed the range of adaptive options later as niches became
12	filled via intra-clade speciation, various additional factors could have shaped subsequent
13	stasis, including stabilizing selection, genetic constraints owing to pleiotropy,
14	developmental or functional constraints and the enduring isolation of the continent
15	which lacked a diversity of other arboreal competitors, among others (Gavrilets and
16	Losos, 2009; Wiens et al., 2010).
17	

18 *4.3. The timing of platyrrhine diversification*

Although obtaining absolute dates for the platyrrhine tree was not the main focus of this report since most of the analyses conducted here are only concerned with relative times, having an accurate estimation of the time of origin of the extant taxa is an important step for incorporating the paleontological information. In a recently published article (Perez et al., 2013), we used different approaches to confidently estimate the absolute time of origin of extant platyrrhines and their main lineages. Results showed

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1	that the most recent common ancestor of extant lineages probably existed between 21-
2	29 to 27-31 Ma, according to two different methodologies, thus indicating that the
3	oldest known Patagonian fossils fall well inside the age range of crown platyrrhines.
4	These dates are broadly concordant with previous works (<i>e.g.</i> , Opazo et al., 2006;
5	Wilkinson et al., 2011). Particularly, if we accept these dates for the initial branching of
6	extant platyrrhines, we could relate the shape of extant phylogeny and the lineages
7	diversification results to the extinction and paleoenvironmental changes in South-
8	America after the Middle Miocene, as is suggested by the analyses of extant and fossil
9	platyrrhine lineages (Fig. 2; Hoorn et al., 2010).
10	Although the question of the Patagonian forms being extinct members of extant
11	lineages or representatives of a separate radiation cannot be resolved only with dating
12	approaches, the absolute dates obtained in Perez et al. (2013), and other previous works,
13	allow for the first hypothesis to be plausible, thus linking the extant species tree with the
14	fossils' temporal distribution. It has to be noted though, that other available divergence
15	time estimations (e.g., Hodgson et al., 2009; Chiou et al., 2011) support a different
16	interpretation of the relationship of the oldest fossil species with extant lineages (but see
17	Perez et al. [2013] for a discussion). In this latter case our results about the shape of
18	platyrrhine lineages and phenotypic diversification are still valid.
19	

20 **5.** Conclusions

The temporal pattern of lineage accumulation and the mode of phenotypic evolution described here based on the extant platyrrhine species might seem contradictory since, as described above, one of the adaptive radiation scenarios predicts

24 an early burst of species origination accompanied by a marked phenotypic

1	diversification. Although both processes may be unlinked, when we also consider the
2	fossil record information this contradiction diminishes as a pattern of an early burst of
3	species diversification arises. This result also shows how the signature of an adaptive
4	radiation may have been erased from the phylogenetic structure of extant species —
5	perhaps by high rate of extinction among lineages after the earlier diversification—but
6	be still retained in the patterns of phenotypic variation, as recent examples have
7	suggested (e.g., Slater et al., 2010; Derryberry et al., 2011). Furthermore, body mass
8	niches appear to have been filled very early in the history of the clade, with a pattern
9	that is remarkably stronger than that seen in other groups. This point also is confirmed
10	by the inclusion of information from the fossil record. Therefore, using a novel
11	approach, as well as fossil and extant species, our analyses support the hypothesis that
12	the platyrrhine tempo and mode of diversification appears to be characterized by an
13	early and profound differentiation in body size related to a multidimensional niche
14	model, followed by little subsequent change (i.e., stasis) in body size. It also suggests an
15	early high rate of extinction among lineages within crown Platyrrhini.

16

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14	Figure Legends
15	Figure 1. Maximum Clade Credibility (MCC) chronophylogenetic tree from the
16	BEAST analysis for 78 species of platyrrhine monkeys. Blue bars on the tree indicate
17	95% confidence intervals for estimated node ages. Support is indicated for nodes with $<$
18	0.8 posterior Bayesian probability. Shaded area shows estimated temporal range for
19	Patagonian fossil lineages.
20	
21	Figure 2 . Semi-logarithmic plots of lineage accumulation through time (LTT plot). Red
22	bold solid line: LTT plot based on the MCC tree of extant species. Dark green/light gray
23	solid and thin dashed lines: median and 95% confidence intervals for the expected

24 distribution under a PB or BD model, respectively, simulated using the maximum

1	likelihood rates estimated from the MCC tree. Extinct lineages are included in the BD			
2	curve. Blue dashed line: observed fossil-based lineage origination curve. The number of			
3	species in extinct taxa was estimated by applying a conservative mean species/genus			
4	ratio of the living taxa (i.e., five) (Quental and Marshal, 2010). Difference between the			
5	area under the MCC tree/Fossil curve and BD curve = 5.05/30.68 ; PB curve =			
6	0.48/26.11. Area between Fossil and MCC curves = 25.63. For interpretation of the			
7	color references in this figure, the reader is referred to the online version.			
8	5			
9	Figure 3. Relative disparity-through-time plot (DTT) for body mass in the 78			
10	platyrrhine species. Solid red line shows disparity calculated for the MCC platyrrhine			
11	tree. Black dashed line shows median expected disparity under the null hypothesis of			
12	random evolution of body size (data generated by 1000 simulations). Shaded area shows			
13	the 95% range for the simulations. Dark gray lines: disparity for 100 trees randomly			
14	sampled from the post-burn in fraction of the BEAST analysis. For interpretation of the			
15	color references in this figure, the reader is referred to the online version.			
16				
17	Figure 4. Adaptive regime models for evolution of body mass (OU models). Each			
18	species was assigned a niche based on a Principal Component analysis (PC, center			
19	figures) according to different hypotheses. Axes represent the principal component			
20	scores used to reduce the number of variables and avoid multicollinearity. Diet			
21	composition niche hypotheses: based on Norconk et al. (2009) diet data. Diet quality			
22	niche hypotheses: based on Allen and Kay et al. (2011). Locomotion niche hypothesis:			
23	based on Youlatos and Meldrum (2011). Multidimensional niche hypothesis: based on			
24	Rosenberger (1992). Note that this hypothesis is not based on a PC analysis.			

1 Ancestral states in the trees were reconstructed using a maximum likelihood criterion.

- 2 See text for details on each model. For interpretation of the color references in this
- 3 figure, the reader is referred to the online version.

- 4
- 5 **Figure 5**. Log body mass for 78 extant species and fossil genera of platyrrhine monkeys
- 6 following the multidimensional niche hypothesis (Rosenberger, 1992; OU-MD5, see
- 7 text and Fig. 4). Fossil genera were assigned to each category according to proposed
- 8 phylogenetic relationships based on morphology (Tejedor, 2008, 2012; Rosenberger et
- 9 al., 2009); however, the phylogenetic relationships of Patagonia fossils are strongly
- 10 discussed (Kay et al., 2008). Circles: La Venta fossils; asterisks: Patagonian fossils.
- 11
- 12











1 Table 1. Mean parameter estimates and comparison of the fit of different lineage

- 2 diversification models to 100 trees randomly drawn from the post-burn in fraction of the
- 3 BEAST analysis (numbers in parentheses are standard deviations; SD). Ma: mega
- 4 annum. r: net diversification rate (speciation extinction; per million years); a: turnover
- 5 (extinction/speciation). Rates go from past to present. AIC: Akaike Information
- 6 Criterion score; wAIC: akaike weights. For the BDS analysis, the best fitting model is
- 7 given. *calculated only for the MCC tree.

Model	Rate shift times (Ma)	r	a	AIC	wAIC
Pure-Birth (2 rates)	0.42 (0.16)	0.17 (0.01); 0.04 (0.05)		-90.03 (9.26)	0.516
BDS (1 shift)	0.42 (0.16)	0.12 (0.01); 0.04 (0.05)	0.44 (0.13); 0.15 (0.16)	-88.75	0.273
Pure-Birth (1 rate)		0.16 (0.01)	-	-86.55 (8.69)	0.091
Birth-Death		0.14 (0.01)	0.22 (0.11)	-85.11 (8.83)	0.044
DDX	-	0.12 (0.02)	-	-85.05 (8.82)	0.043
DDL	-	0.16 (0.03)	-	-84.56 (8.69)	0.034
DDD+E*	-	0.52	0.34	-69.94	0.000

8 9

1 Table 2. Performance of alternative models of body mass evolution in the platyrrhine 2 diversification. Models were fit over 100 trees randomly drawn from the post burn-in 3 fraction of the BEAST analysis. Refer to text for description of models. AICc: Median 4 and standard deviation (SD, in parentheses) for Akaike Information Criterion scores 5 corrected for sample size. Δ AICc: difference between each model median AICc score 6 and the best fitting model AICc score; wAICc: weighted AICc. α : median and SD for

7	the strength	of selection	parameter	estimation	(see methods).
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Model	AICc	ΔAICc	wAICc	α
OU-MD5	488.23 (9.58)	0	0.9988	1.91 (1.06)
OU-Loc3	503.48 (13.17)	14.24	0.0008	0.27 (0.24)
OU-DietC4	504.61 (13.44)	15.37	0.0004	0.23 (0.26)
BM	509.48 (13.48)	20.24	0	-
OU-DietC3	512.97 (12.82)	23.73	0	0.12 (0.18)
OU-DietQ4	514.14 (12.77)	24.90	0	0.13 (0.20)
	K			
OU-DietQ5	515.77 (12.85)	26.53	0	0.13 (0.20)
OU1	516.76 (12.78)	27.52	0	0.06 (0.07)

0

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1 Highlights

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- 2 We estimate a dated platyrrhine phylogenetic tree for ca. 80% extant species.
- 3 Extant lineage diversification shows a constant rate of accumulation.
- 4 In contrast, body size diversification shows a pattern of early niche partition 5 followed by stasis.
- 6 Body size evolved according to the occupation of a multidimensional niche.
- 7 When oldest fossils are included, an early pulse of lineage origination appears.
- 8 9