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Unresolved molecular phylogenies of gibbons and siamangs (Family: Hylobatidae) based on mitochondrial, Y-linked, and X-linked loci indicate a rapid Miocene radiation or sudden vicariance event

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ABSTRACT

According to recent taxonomic reclassification, the primate family Hylobatidae contains four genera (Hoolock, Nomascus, Symphalangus, and Hylobates) and between 14 and 18 species, making it by far the most species-rich group of extant hominoids. Known as the ''small apes'', these small arboreal primates are distributed throughout Southeast, South and East Asia. Considerable uncertainty surrounds the phylogeny of extant hylobatids, particularly the relationships among the genera and the species within the Hylobates genus. In this paper we use parsimony, likelihood, and Bayesian methods to analyze a dataset containing nearly 14 kilobase pairs, which includes newly collected sequences from X-linked, Y-linked, and mitochondrial loci together with data from previous mitochondrial studies. Parsimony, likelihood, and Bayesian analyses largely failed to find a significant difference among phylogenies with any of the four genera as the most basal taxon. All analyses, however, support a tree with Hylobates and Symphalangus as most closely related genera. One strongly supported phylogenetic result within the Hylobates genus is that Hylobates pileatus is the most basal taxon. Multiple analyses failed to find significant support for any singular genus-level phylogeny. While it is natural to suspect that there might not be sufficient data for phylogenetic resolution (whenever that situation occurs), an alternative hypothesis relating to the nature of gibbon speciation exists. This lack of resolution may be the result of a rapid radiation or a sudden vicariance event of the hylobatid genera, and it is likely that a similarly rapid radiation occurred within the Hylobates genus. Additional molecular and paleontological evidence are necessary to better test among these, and other, hypotheses of hylobatid evolution.

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1. Introduction

Gibbons and siamangs are hominoid primates of the family Hylobatidae, which is the sister group to Hominidae, the group that includes humans. According to recent taxonomic reclassification, there are between 14 and 18 species of extant hylobatids [\(IUCN,](#page-7-0) [2010; Mootnick, 2006; Thinh et al., 2010](#page-7-0)), making this the most species-rich group of extant apes. Inhabitants of the rainforests of Southeast, South and East Asia, all hylobatid species are listed on the IUCN Red List of Threatened Species, with most of them

endangered ([IUCN, 2010](#page-7-0)). An accurate phylogeny of these apes would enhance our understanding of Asian biogeography, morphological evolution, hominoid paleontology, and contribute to conservation efforts.

Multiple different data types have been studied to address hylobatid phylogenetics, including morphological, vocal, karyotypic, and molecular genetic data [\(Chatterjee, 2006; Groves, 1972;](#page-7-0) [Haimoff et al., 1982; Hayashi et al., 1995; Matsudaira and Ishida,](#page-7-0) [2010; Muller et al., 2003; Roos and Geissmann, 2001; Takacs et al.,](#page-7-0) [2005; Thinh et al., 2010; Whittaker et al., 2007](#page-7-0)). One point of agreement among molecular studies is that hylobatids can be represented by four monophyletic genera: Hylobates, Symphalangus, Nomascus, and Hoolock [\(Brandon-Jones et al., 2004\)](#page-7-0). However, genetic studies conflict on two key aspects of hylobatid phylogeny. First, there is disagreement on the relationships among the four genera. Several genetic studies forward Nomascus as the most basal taxon ([Chatterjee, 2006; Roos and Geissmann, 2001; Thinh et al., 2010\)](#page-7-0),

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while others, including those based on chromosomal data, give support for basal positioning of Hoolock [\(Muller et al., 2003; Takacs et al.,](#page-7-0) [2005\)](#page-7-0). Morphological evidence has supported a basal position for Symphalangus [\(Groves, 1972](#page-7-0)). Secondly, within Hylobates there is considerable uncertainty regarding the relationships among the different species. One important question relates to the position of the most basal member of Hylobates. Hylobates pileatus has been suggested to be the most basal member in studies based on molecular and morphological data [\(Geissmann, 2002; Takacs et al., 2005;](#page-7-0) [Whittaker et al., 2007](#page-7-0)), while other analyses support a Hylobates klossii as the basal member of Hylobates (e.g. [Purvis, 1995\)](#page-7-0).

Using molecular phylogenies, there have been different interpretations of the hylobatid evolutionary history. [Chatterjee](#page-7-0) [\(2006\)](#page-7-0) forwarded a scenario where hylobatids began their radiation in Eastern Indochina at about 10.5 Ma and subsequent waves of radiation occurred predominantly southwardly, with a westward radiation of Hoolock [\(Chatterjee, 2006\)](#page-7-0). [Whittaker et al.](#page-8-0) [\(2007\)](#page-8-0) examined Hylobates and suggested a general north to south biogeographic expansion. [Thinh et al. \(2010\)](#page-8-0) suggested that the intergeneric hylobatid radiation occurred on the Southeast Asian mainland in the Miocene, specifically in the area of the Hengduan Mountains. Subsequently, the genera migrated into their present positions on the mainland. This was followed by a southward expansion by Hylobates onto the Sunda shelf, where vicariant speciation mainly occurred.

In the present study, we collected nuclear DNA sequences from the X-linked G6PD locus and the Y-linked ZFY locus, and the mitochondrial loci COII and ND4/5 from a number of hylobatids. These sequences were added to a comprehensive dataset of mitochondrial DNA sequences, including five complete mtDNA genomes ([Matsudaira and Ishida, 2010](#page-7-0)). This alignment was analyzed using parsimony, likelihood, and Bayesian phylogenetic methods, and multiple hypotheses of hylobatid phylogeny were tested. A molecular clock analysis was used to estimate the date of speciation events in the evolutionary history of hylobatids. The results are discussed within the context of recently proposed hypotheses of hylobatid evolutionary history.

2. Materials and methods

2.1. Collection of DNA sequences

Gibbon and outgroup samples were provided by several individuals and institutions. A list of samples, including identification

Table 1

Sample information, including sequenced regions.

number, institution, and genes for which sequences are included in this study is shown in Table 1. Many different sample types were used for this study: tissue, whole blood, serum, hair, urine, and tissue culture cell suspensions. Extraction of all samples was performed using the IsoQuick DNA Extraction Kit (ORCA Research; Bothell, WA), a guanidine thiocyanate-based method, according to the manufacturer's protocols.

DNA sequence data were collected from two mitochondrial regions, one X-linked locus, and one Y-linked locus. The first is the 684 bp (base pair) cytochrome oxidase subunit II locus (COII). The second is an 898 bp fragment consisting of sequences from two NADH dehydrogenase complex genes (ND4 and ND5). The third data set is from four introns of the X-linked Glucose-6-phosphate dehydrogenase gene (introns 4, 5, 7, and 8, also called D, E, G, & H). The final data set is from the last intron of the Y-linked zinc finger gene (ZFY). Polymerase chain reaction (PCR) was used to amplify target DNA for both regions in preparation for sequencing. For COII, the entire coding sequence was amplified with primers listed in Supplementary materials. For the ND4/5 region, primers A–F, described by [Hayashi et al. \(1995\)](#page-7-0), were used. Because of the possibility of introgression of mitochondrial sequences into the nucleus, multiple independent PCRs with different primer combinations were performed for all samples. Sequences from all amplifications matched exactly, indicating that each of them came from a single population of target DNA. None of the consensus sequences contained stop codons or other anomalous codons, and each was therefore considered to represent the true mitochondrial sequence. For G6PD, amplification was done using primers located within the flanking exons. In some occasions, nested PCR was used. For ZFY, amplification was done using primers located within the flanking exons using nested PCR.

PCR conditions were as follows: up to 10 µL of DNA (concentration unquantified, from 50 µL of IsoQuick extracted DNA) was used as a template in a 50 µL reaction containing 10 mM Tris-HCl pH 8.3, 50 mM KCl, 20–25 pM of each primer, 1.25–2.5 units of polymerase (AmpliTaq, Perkin-Elmer/ABI, Foster City, CA, or Qiagen Taq, Valencia, CA), $1.25-1.5$ mM MgCl₂, and 0.15-0.2 mM of each dNTP. Reactions were run in either a MJ Research Minicycler or a Perkin-Elmer 9600 thermocycler. Cycling parameters for hylobatids were as follows:

For COII a 3 min 94 \degree C denaturation was followed by 30 repetitions of a 60 s 94 °C, 60 s 48 °C, 60 s 72 °C cycle. For $ND4/5$ a 3 min 94 °C denaturation was followed by 30 repetitions of a 45 s 94 °C, 60 s 48 °C, 60 s 72 °C cycle. For G6PD gibbon samples, a 3 min 94 °C denaturation was followed by 40 repetitions of a 30 s 94 °C,

45 s 65 °C, 60 s 72 °C cycle. For the non-gibbon hominoids, the cycle consisted of 30–35 repetitions of 60 s at 94 °C, 80 s at 65 °C, and 130 s at 72 °C. For introns D and E, 4 s were added to the extension with each cycle. For Papio and Macaca, the cycle consisted of 30–35 repetitions of 30 s at 93 °C, 30 s at 60–64 °C, and 45 s at 72 °C. For ZFY a 3 min 94 \degree C denaturation was followed by 40 repetitions of a 30 s 94 °C, 45 s 50 °C, 45 s 72 °C cycle. A final cycle of 5-10 m at 72 °C cycle followed.

Fragments were gel-purified and used directly in sequencing reactions. DNA sequencing of double-stranded PCR product was performed using the ABI Prism Dye-Deoxy reaction kit (Applied Biosystems Inc., Foster City CA) according to the manufacturer's protocol. Unincorporated primers and nucleotides were removed using Centriflex bead matrix columns (Edge BioSystems, Gaithersburg, MD) according to the manufacturer's protocols. Column eluant was dried down in a Speed-Vac with no heat, resuspended in 3.5μ L of loading buffer (Formamide, 50 mM EDTA, 30 mg/ml dextran blue) and run on a 4.75 Sequagel (National Diagnostics) or a 6% Long Ranger acrylamide gel (FMC, Rockland, ME) for 12 h on an ABI 373 automated sequencer.

2.2. Alignment of DNA sequences

The DNA sequences collected from gibbons were added to the available DNA sequences of hylobatids, Homo sapiens, Pan troglodytes, Pan paniscus, Gorilla gorilla, Pongo abelii, Pongo pygmaeus, Macaca mulatta, and Papio from Genbank and the literature (Table 2). This included five complete hylobatid mitochondrial genomes (Hylobates agilis, Hylobates pileatus, Symphalangus syndactylus, Nomascus siki, and Hylobates lar) and additional species for the mtDNA loci cytochrome b (CytB), cytochrome oxidase II (COII), and

Table 2

the NADH dehydrogenases (ND3, ND4, ND4L, and ND5) (Table 2). Only the protein coding regions of the mtDNA genomes were used because the non-protein coding regions were not straightforward to align. The protein coding genes were aligned using ClustalW ([Thompson et al., 1994\)](#page-8-0), separated by gaps in order to preserve the reading frame.

2.3. Phylogenetic analyses

In the phylogenetic analyses each gene was analyzed separately as well as in a single combined multi-gene alignment of all data. For maximum likelihood and Bayesian analyses, this alignment was analyzed as five independent data partitions (mtDNA first, second, and third codon positions, G6PD, and ZFY). This method was chosen based on the results of a recent study ([Ren et al., 2009\)](#page-7-0) which showed that likelihood performed better than (i) a 'supermatrix' method that concatenated partitions into a single alignment analyzed using one set of parameters and (ii) a 'supertree' method that separated partitions, estimated a phylogenetic tree for each partition, and used supertree methods to combine independent trees [\(Ren et al., 2009](#page-7-0)). In each analysis, Papio and/or Macaca were used as the outgroup taxa to the ingroup, which included both hylobatids and the other hominoids, Homo, Pan, Pongo, and Gorilla.

A parsimony-based phylogenetic analysis was conducted in PAUP^{*} v. 4.0b10 [\(Swofford, 2001](#page-8-0)). Again, the three separate loci were analyzed both independently and together. In all cases, two separate character-weighting matrices were used. For mtDNA, a 10:1 weighting scheme was used, with transversions being weighed more heavily than transitions. For G6PD and ZFY, a 2:1 weighting scheme was used, with transversions being weighed more heavily than transitions. These weightings reflect the greater

frequency of transitions than transversions in molecular data, with an increased bias in mitochondrial DNA. Papio and Macaca were used as outgroups, and a branch-and-bound tree search was performed to find the most parsimonious tree(s). A parsimony-based bootstrap analysis was conducted using 1000 search replicates, using a heuristic search at each replicate of 10 random taxon additions. For the combined alignment, phylogenetic hypothesis testing was done using constraint trees with each genus as basal and subsequently conducting a heuristic search. These constrained trees were compared to the most parsimonious trees using the [Templeton \(1983\)](#page-8-0) and winning sites tests [\(Prager and Wilson,](#page-7-0) [1988\)](#page-7-0) using PAUP*.

A maximum likelihood-based phylogenetic analysis was conducted in RAxML v. 7.0.4 [\(Stamatakis, 2006; Stamatakis et al.,](#page-8-0) [2008\)](#page-8-0). G6PD, ZFY, and mitochondrial loci were analyzed separately, with the mtDNA partition having three different partitions corresponding to each codon position. A combined alignment of all data using five separate partitions was also analyzed. For all partitions a general time reversible model of evolution with a gamma model for among site rate variation was used (GTRGAMMA), and Papio and Macaca were used as outgroups in all analyses. A maximum likelihood bootstrap analysis with 1000 replicates was also conducted on the combined alignment. Phylogenetic hypothesis testing was done on the combined alignment using constraint trees with each genus as basal and subsequently conducting a maximum likelihood search. These constrained trees were compared to the most likely tree using the SH test ([Shimodaira and Hasegawa,](#page-8-0) [1999\)](#page-8-0) in RAxML.

A Bayesian-based phylogenetic analysis was conducted with an MPI enabled MrBayes v. 3.1.2 ([Altekar et al., 2004; Huelsenbeck](#page-7-0) [and Ronquist, 2001; Ronquist and Huelsenbeck, 2003](#page-7-0)). Three separate partitions were used for the mtDNA sequence data, each with a general time reversible model of evolution with six substitution classes (Nst = 6) and a Γ and invariant (Invgamma) model for among site rate variation. For the G6PD data, a model with two substitution classes (Nst = 2) and a gamma model for among site rate variation was used. For the ZFY data, a model with six substitution classes and a gamma model for among site rate variation was used. A Dirichlet prior of 1 was used for the each of the base state frequencies in each of the four partitions. These models were chosen using MrModelTest v. 2 ([Nylander, 2004](#page-7-0)). The site-specific rate model was set to variable, allowing the rate to vary across partitions. All of these parameters were unlinked across the partitions. Macaca mulatta was used as the outgroup (only one outgroup taxon is allowed by MrBayes). Three independent runs were initiated each using a random starting tree; each run consisted of five million steps, with sampling every 100 steps, and used four chains and a heating parameter of 0.085, allowing for good swapping between chains. The first 20% of each run was discarded as burnin. Convergence was assessed among chains by examining the potential scale reduction factors (PSRF), effective sample sizes (ESS) (estimated by Tracer v. 1.4.1 [\(Rambaut and Drummond, 2007\)](#page-7-0)), and the similarity of parameter estimates among runs. A similar analysis was conducted for each independent locus as well. Phylogenetic hypothesis testing was done on the combined alignment using constraint trees with each genus as basal and subsequently conducting a Bayesian analysis. The harmonic mean likelihoods of the constrained trees were compared to the unconstrained trees in a manner similar to [Nylander et al. \(2004\)](#page-7-0).

2.4. Molecular clock analysis

A molecular clock analysis was conducted using the Bayesian MCMC program mcmctree of PAML v. 4.3 ([Yang, 2007\)](#page-8-0) using a user-input tree matching the Bayesian consensus tree ([Fig. 3\)](#page-5-0). This tree was chosen because a completely bifurcating tree is required.

The alignment was broken into four separate partitions and analyzed together using the HKY + Γ 5 model [\(Hasegawa et al., 1985;](#page-7-0) [Yang, 1994](#page-7-0)), a model with different transition/transversion rate ratios (κ) , different base frequencies and different gamma shape parameter α for each of the four partitions. Gamma priors were assigned on parameters $\kappa \sim G(6, 2)$, with mean 3, and $\alpha \sim G(1, 1)$. Substitution rates are assumed to drift over time independently among the partitions. A 100 Myr time unit was used. The rate at the root is the gamma prior $\mu \sim G(1, 0.4)$. A geometric Brownian motion model was used to accommodate substitution rate variation across lineages ([Rannala and Yang, 2007\)](#page-7-0). The rate-drift parameter was assigned the gamma prior $\sigma^2 \sim G(1, 10)$. The prior of times was generated from the birth to death process with sampling, with parameters $\lambda = \mu = 1$ and $\rho = 0$, so that the kernel is uniform [\(Yang and Rannala, 2006](#page-8-0)). Two independent runs were conducted, each with a 10,000 step burnin followed by 100,000 steps, with sampling every five steps. Convergence was assessed by agreement between two independent runs. The tree was calibrated by assigning divergence times to two independent nodes. A divergence time of 6–8 Ma for the Homo/Pan split was based on the presence of the candidate hominins Orrorin tugenensis $(\sim$ 5.8 Ma ago) ([Senut et al., 2001](#page-7-0)), Ardipithecus kadabba (5.2– 5.8 Ma ago) [\(Haile-Selassie et al., 2004\)](#page-7-0), and Sahelanthropus tchadensis (6.8–7.2 Ma ago) [\(Lebatard et al., 2008](#page-7-0)). A divergence time of 6–8 Ma ago was also assumed for the Macaca/Papio split based on the age of the earliest members of Macaca (5.5 Ma) and paleontological estimates for the divergence of these taxa between 7 and 8 Ma ago ([Delson, 1992, 2000\)](#page-7-0). The root node of the tree (crown catarrhine) was calibrated using a gamma distribution with a shape parameter of 110 and scale parameter of 1/400. This corresponds to a distribution with the 2.5% date at 22.6 Ma, 97.5% date at 32.9 Ma, and mean date of 27.4 Ma. This is based on the date of the earliest crown catarrhine, the hominoid Morotopithecus, from 20.6 Ma ago [\(Gebo et al., 1997](#page-7-0)). This distribution generates an upper bound, bracketing a conservatively large estimate for the origin of crown catarrhines. Furthermore, these calibrations are 'soft' bounds ([Yang and Rannala, 2006\)](#page-8-0), i.e. the date estimates for these nodes have a limited prior probability (2.5%) to be older or younger than the calibrations.

3. Results

The final alignment of mitochondrial coding sequences totaled 10,854 base pairs, the final G6PD alignment totaled 2191 base pairs, and the ZFY alignment totaled 825 base pairs. Using all analytical methods, the single locus analyses offered little phylogenetic resolution (see Supplementary materials). For example, in analyses of mtDNA sequences, the position of Hoolock differed among the different optimality criteria; in analyses of the G6PD locus the positions of the genera relative to one another was different among optimality criteria; and in ZFY the limited differences among the species caused little resolution among hylobatids and long branch attraction between the hylobatid and Pongo lineages. Because of the lack of resolution of the single gene analyses, the results from the combined analysis are presented in greater depth.

3.1. Parsimony results

The maximum parsimony analysis resulted in two most parsimonious trees. Both trees agreed in their placement of Nomascus as the most basal taxon, with Hoolock as next most basal, then a Symphalangus + Hylobates clade. Within the Hylobates genus, the two most parsimonious trees differed in their placement of Hylobates agilis. The bootstrap value for the Hoolock node was 52% and for the Symphalangus + Hylobates node was 65%. A bootstrap

consensus tree appears in Fig. 1. There was strong support for Hylobates as a monophyletic group (100%) and for the placement of Hylobates pileatus as the most basal member of this group. Within the remaining Hylobates species, however, the bootstrap values on all the branches were low (27–60%).

3.2. Likelihood results

The maximum likelihood tree differed from the most parsimonious trees in the placement of the most basal taxon (Fig. 2). In the maximum likelihood tree, Hoolock is the most basal genus, Nomascus is the next most basal, followed by a Symphalangus + Hylobates clade. Within the Hylobates genus, H. pileatus is most basal, but the placement of the remaining species differs from both most parsimonious trees. Bootstrap support for the placement of Hoolock was very low (33%). The placement of Symphalangus was moderately supported (80%). Within Hylobates, the bootstrap values varied.

3.3. Bayesian results

The Bayesian analysis resulted in a tree similar to the maximum likelihood tree at the genus level ([Fig. 3\)](#page-5-0). Hoolock was placed as the most basal genus. Nomascus was the next most basal, with a clade credibility value of 61. Symphalangus was sister taxon to Hylobates with a clade credibility value of 85. Within Hylobates, Hylobates pileatus was the most basal species. The placement of the remaining species in the Hylobates genus differed from both the maximum parsimony trees and the most likely tree.

Fig. 2. Bootstrap consensus phylogram using maximum likelihood. Bootstrap values appear above branches leading to nodes except within most Hylobates nodes, where they appear to the right of the node. A cladogram that has collapsed the branches of <50% bootsrap support appears in Supplementary materials Fig. S4.

3.4. Hypothesis testing

Because one of the most critical phylogenetic questions assessed in this study is the placement of the most basal taxon, we conducted further testing among hypotheses placing each of the four genera as the most basal hylobatid. A comparison of the two maximum parsimony trees (placing Nomascus basal) against a tree with Hoolock as most basal showed no statistically significant difference between them ([Table 3\)](#page-5-0). Trees placing Symphalangus or Hylobates as most basal were significantly worse than the most parsimonious trees with Nomascus as basal. The likelihood-based SH test found that testing the ML tree against trees with the other genera as basal did not reveal statistically significantly differences among the trees. Similarly, a Bayes factor analysis found no significant differences between trees with Hoolock, Nomascus, Symphalangus or Hylobates as the most basal taxon. These tests show that any resolution among these genera is not statistically robust.

3.5. Molecular clock results

A molecular clock was used to estimate divergence dates based using the Bayesian tree. This tree was chosen because a single completely bifurcating tree is required by the methodology and the Bayesian method yielded relatively high posterior probabilities for the placement of the different genera. The molecular clock analyses yielded a date for the origin of Hylobatidae at 21.8 Ma (19.7–24.1 Ma) and for the radiation of the family at 7.3 Ma (6.4– 8.0 Ma) ([Fig. 4\)](#page-6-0). The subsequent radiation of Nomascus and

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Hylobates moloch 0.99 *Hylobates pileatus Symphalangus syndactylus Nomascus siki Nomascus leucogenys Nomascus gabriellae Hoolock leuconedys* ¹ *Homo sapiens* 1 *Pan troglodytes* 1 -*Pan paniscus* $\overline{1}$ *Gorilla gorilla* 1 *Pongo pygmaeus*

1

Pongo abelii

Fig. 3. Phylogram estimated using Bayesian phylogenetics. Posterior probability values appear above branches leading to nodes except within most Hylobates nodes, where they appear to the right of the node.

Symphalangus + Hylobates followed shortly thereafter (7.0 Ma and 6.4 Ma). The radiation of Hylobates occurred at 3.5 Ma (3.1– 4.0 Ma) (additional dates and parameters are given in Supplementary materials). Dates estimated using trees with the other genera as basal gave qualitatively similar results.

4. Discussion

Combined analyses of the loci offered better phylogenetic resolution among the hylobatids than individual analyses of each gene. While combined analyses of mitochondrial, X-linked, and Y-linked DNA sequence data confirm the existence of the four separate genera of hylobatids, the analyses conflict regarding the relationships among these genera. Parsimony analysis supports Nomascus as the most basal taxon, and some molecular studies have supported this arrangement ([Chatterjee, 2006; Roos and Geissmann, 2001; Thinh](#page-7-0) [et al., 2010\)](#page-7-0). Maximum likelihood and Bayesian analyses support Hoolock as the most basal, and both molecular and karyological

Results of the phylogenetic hypothesis tests.

studies have supported this alternative ([Muller et al., 2003; Takacs](#page-7-0) [et al., 2005](#page-7-0)). Bootstrap values showed moderate support for the particular topology among the hylobatid genera produced by each method. However, hypothesis testing among basal positions for the different genera found few significant differences among these hypotheses. Only a basal position for Symphalangus and Hylobates was rejected using parsimony-based tests; none of the alternatives were rejected using Bayesian- and likelihood-based tests. Within Hylobates, there was some consensus, in that Hylobates pileatus represented the most basal taxon and that Hylobates klossii and Hylobates moloch are sister taxa. The placement of these Hylobates species agrees with other studies ([Geissmann, 2002; Takacs et al.,](#page-7-0) [2005; Whittaker et al., 2007\)](#page-7-0). For the other remaining species within Hylobates, there is no consensus. For example, the placement of Hylobates lar differed in each phylogenetic reconstruction.

Although the present study suggests more phylogenetic uncertainty than suggested by other recent analyses (e.g. [Matsudaira](#page-7-0) [and Ishida, 2010\)](#page-7-0), it represents a significant advance for three reasons. First, the present dataset is based on a large sample of hylobatids, including members of all four genera. Second, the present dataset is the only study that examined regions of three independent genomic compartments. Others have been solely based on mitochondrial sequences. Third, in addition to the phylogenetic analyses, this study included explicit phylogenetic hypothesis testing using parsimony, likelihood, and Bayesian methods.

The persistent phylogenetic uncertainty among hylobatid genera and within Hylobates has a number of potential causes. First, the molecular clock results indicate that the radiation of the genera happened over an interval of less than 1 Ma, and this rapidity of the hylobatid radiation would lead to very short internal branches at the base of the hylobatid phylogeny. Members of the Hylobates genus also experienced a rapid radiation. These radiations have been suggested by others [\(Muller et al., 2003; Takacs et al., 2005;](#page-7-0) [Thinh et al., 2010; Whittaker et al., 2007](#page-7-0)). Ancestral polymorphism followed by incomplete lineage sorting tends to create mismatches between gene trees and the species tree to a greater degree when divergence times are shorter and when ancestral population sizes are larger. Rapid radiations may have been coupled with other biogeographic factors including repeated dispersals, repeated forest expansions and contractions, and vicariant speciation, and all of these factors could lead to a complicated and difficult to resolve species phylogeny. Although an accurate phylogeny would assist in the reconstruction of hylobatid biogeographic history and indeed, some have attempted to use gibbon molecular phylogenies in this manner to posit specific dispersals and vicariant speciation events (e.g. [Chatterjee, 2006, 2009; Thinh et al., 2010](#page-7-0)), these methods are completely dependent on having an accurate, well-resolved species tree. Hypothesis tests in the present study show that such a resolved phylogeny is not yet at hand for hylobatids. Therefore, such biogeographic scenarios are premature.

A second scenario that would result in the observed phylogenetic uncertainty is a sudden vicariance model. In this model, the phylogenetic uncertainty among genera of hylobatids is not interpreted as an artifact of ancestral polymorphisms or a complicated dispersal model, but rather is a real signal in its own right, i.e. a hard polyto-

^a Number of tree steps (Templeton test; winning sites test).

b Likelihood values (SH test).

^c Harmonic mean likelihood (Bayes factor comparison).

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Fig. 4. Tree scaled to the molecular clock dates of each node. Numbers at nodes indicate the 95% Bayesian credibility interval from the posterior distribution. Three calibrations nodes are indicated with a 'C = ' within brackets. The dates following the 'C = ' correspond to the lower (2.5%) and upper (97.5%) bounds of the calibration prior distribution. Other parameter estimates given in Supplementary materials. All dates are in Ma.

my. A similar model was proposed for the biogeographic history of Hylobates species [\(Whittaker et al., 2007](#page-8-0)) and the model may also apply to the intergeneric radiation of hylobatids. There is paleoclimatological and paleoecological evidence to support the vicariance model. Here it is assumed that the ancestor of hylobatids was widely distributed throughout the rainforests of Southeast, South, and East Asia during the relatively warm and moist Miocene. From the middle Miocene until about 6 Ma, global climate cooled ([Zachos](#page-8-0) [et al., 2001](#page-8-0)). There is evidence for a habitat shift at this time from a nearby site in the Surai Khola area (13–1 Ma) of the Nepalese Siwaliks [\(Corvinus and Rimal, 2001\)](#page-7-0). From about 13 to 9.5 Ma the landscape was dominated by a tropical evergreen forest type that this similar to today's Southeast Asian rainforests ([Corvinus and](#page-7-0) [Rimal, 2001\)](#page-7-0). Subsequently, from about 9.5 to 7.5 Ma there is evidence for moist deciduous forests at the site. Around the Miocene–Pliocene boundary, the habitat was drier and the subtropical environment was varied, with only about 25% evergreen forests ([Corvinus and Rimal, 2001](#page-7-0)). During this cooling and habitat shift, the rainforest habitat of hylobatids might have fragmented or shrunk relatively simultaneously into multiple discontiguous distributions throughout Southeast, South, and East Asia. This model proposes that a large contiguous hylobatid distribution fragmented simultaneously and vicariantly into four descendant populations that later evolved into the different hylobatid genera. Such a simultaneous vicariant speciation event would likely lead to phylogenetic uncertainty among the hylobatid genera. The estimated timing of generic radiation produced by this study supports this model. The biogeography of other Southeast Asian mammals has similarly been interpreted as the result of vicariant events in the Late Miocene and Pliocene (e.g. [Gorog et al., 2004; Steiper, 2006\)](#page-7-0).

Within Hylobates, a similar model seems apparent. Within this genus, there was agreement among the analyses at two nodes, Hylobates pileatus as the most basal species and for a sister taxon relationship between Hylobates klossii and Hylobates moloch. There was no agreement for the placement of the other three species or the other nodes. The current distribution of Hylobates pileatus suggests that Hylobates may have originated on the mainland while derived members of the genus dispersed west and southward.

Once distributed throughout Southeast Asia, cooling may have caused a second round of in situ vicariant speciation. The timing of these nodes roughly coincides with the onset of the Northern Hemisphere Glaciation at 3.2 Ma ([Zachos et al., 2001](#page-8-0)), potentially linking these phenomena. It is noteworthy that there is some agreement that Hylobates klossii and Hylobates moloch are sister taxa, though with varying degrees of support. Hylobates klossii occurs on the Mentawai Islands, west of Sumatra, while Hylobates moloch occurs on Java, implying a dispersal event between these locations. Overall, this interpretation supports [Whittaker et al.](#page-8-0) [\(2007\)](#page-8-0) and suggests that cooling events may have also driven vicariant speciation within Hylobates.

The two biogeographic models make different predictions for future studies of gibbon molecular phylogenetics. A rapid divergence, multiple dispersal, and vicariance model could have potentially led to ancestral polymorphisms that would cause multiple incongruences between gene trees and species trees. However, this model would predict that a robust phylogeny for hylobatids would emerge with additional study. This situation may parallel the situation with New World primates and African hominoids, taxa that required a considerable amount of data from a number of independent loci to resolve a robust phylogeny (Ray et al., 2005; Ruvolo, 1997; Wildman et al., 2009). In situ vicariance events, on the other hand, makes the prediction that additional analyses of hylobatids will not uncover a robust tree. Instead, this model predicts the intergeneric phylogeny of gibbons and many of the intrageneric nodes within Hylobates will remain resistant to the recovery of an accurate and well-supported bifurcating tree. In either case, examining a large number multiple independent loci will be required to assess hylobatid phylogenetics with confidence. Paleontological evidence is also of importance in testing among these hypotheses. Unfortunately, there is a relative lack of hylobatid fossils from critical time periods and localities. Additional paleontological research throughout Southeast, South and East Asia at Miocene sites that are inferred to be rainforest would greatly improve our understanding of hylobatid evolutionary history.

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at [doi:10.1016/j.ympev.2010.11.005.](http://dx.doi.org/10.1016/j.ympev.2010.11.005)

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