Molecular Phylogeny of the Bamboo Sharks (Chiloscyllium spp.)

Noor Haslina Masstor, Abdullah Samat, Shukor Md Nor, and Badrul Munir Md-Zain

School of Environmental and Natural Resource Sciences, Faculty of Science and Technology, Universiti Kebangsaan Malaysia, 43600 Bangi, Selangor, Malaysia

Correspondence should be addressed to Badrul Munir Md-Zain; abgbadd1966@yahoo.com

Received 27 February 2014; Revised 1 May 2014; Accepted 16 May 2014; Published 11 June 2014

Academic Editor: Barbara Mantovani

Copyright © 2014 Noor Haslina Masstor et al. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Chiloscyllium, commonly called bamboo shark, can be found inhabiting the waters of the Indo-West Pacific around East Asian countries such as Malaysia, Myanmar, Thailand, Singapore, and Indonesia. The International Union for Conservation of Nature (IUCN) Red List has categorized them as nearly threatened sharks out of their declining population status due to overexploitation. A molecular study was carried out to portray the systematic relationships within Chiloscyllium species using 12S rRNA and cytochrome b gene sequences. Maximum parsimony and Bayesian were used to reconstruct their phylogeny trees. A total of 381 bp sequences’ lengths were successfully aligned in the 12S rRNA region, with 41 bp sites being parsimony-informative. In the cytochrome b region, a total of 1120 bp sites were aligned, with 352 parsimony-informative characters. All analyses yield phylogeny trees on which C. indicum has closer relationships with C. plagiosum. C. punctatum is sister taxa to both C. indicum and C. plagiosum while C. griseum and C. hasseltii formed their own clade as sister taxa. These Chiloscyllium classifications can be supported by some morphological characters (lateral dermal ridges on the body, coloring patterns, and appearance of hypobranchials and basibranchial plate) that can clearly be used to differentiate each species.

1. Introduction

Over 450 sharks species with eight orders have been recognized. Chiloscyllium, within the family Hemiscylliidae, order Orectolobiformes, is one of the disperse populations in the ocean as their habitat extend from Indian Ocean to Western Pacific. There are seven identified species, of which only five living species have been recorded inhabiting the ocean of Malaysia Chiloscyllium griseum [1], C. indicum [2], C. punctatum [1], C. plagiosum [2], and C. hasseltii [3] by the Department of Fisheries Malaysia [4]. For Chiloscyllium, their hardiness, small size, and beautiful patterning have caused them to be commonly imported for the pet trade [5]. C. griseum, C. plagiosum, and C. punctatum particularly are among the popular aquarium species and being kept in public aquaria [6]. Therefore, for a long time, sharks were overhunted, resulting in their globally declining populations in spite of populating wide range of water areas. Sharks are susceptible to overfishing and slow to recover if overfished, due to their low reproductive rate, late maturity, and small population size. To overcome this challenge, many conservation efforts have been introduced and applied, including listing them as IUCN Red List status. Listed as “near threatened” by the IUCN Red List [7], our study interest, Chiloscyllium species, has also faced decreasing population problems.

Species identification, taxonomy, and molecular systematics of shark species are among the initial steps in conserving and managing shark stocks. In spite of that, sharks phylogenetic relationships are still not well established even after 150 years of study as reported by Naylor et al. [8]. Early interest in the systematics among elasmobranchs had led to the descriptions of families of sharks and rays. Müller and Henle [9] study had established the genus Chiloscyllium. In 1841, C. indicum and two new species, C. griseum and C. punctatum, were assigned to the genus of Chiloscyllium. Bleeker [3] described C. hasseltii, which for a long time was considered a synonym of C. griseum. Waite [10] first placed genus Hemiscyllium and genus Chiloscyllium in their own family, Hemiscylliidae. However, Regan [11] assigned all Chiloscyllium species to the family Orectolobidae. Whitley [12] placed Hemiscyllium and Chiloscyllium back in family...
Hemiscylliidae, and this was maintained by Compagno [13]. Dingerkus and DeFino [5] identified and described two new *Chiloscyllium* species, *C. burmensis* and *C. arabicum*. Unfortunately, all of those studies were only based on the analysis of morphological data, including anatomical differences such as the neurocranium, dorsal and anal fins, visceral arch, shoulder girdle and pectoral fin, pelvic girdle and clasper, vertebræ and caudal fin, head sensory canal system, and other external morphologies [14], skeletal anatomy and dermal denticles [5], and diagnostic features and biology [6]. Those studies made a great impact on the classification of this genus, but, with the advent of molecular techniques, it can provide supportive information in various fields of study such as the molecular systematics, biogeography, population genetics, and evolutionary history [15, 16]. There are some numbers of molecular studies that had been conducted to review the phylogeny of sharks at the order and family levels [8, 17, 18] and at the species level by Human et al. [19].

Even though there were some efforts to clarify shark species relationships, as stated by Naylor et al. [8] and Heinicke et al. [20] the phylogenetic relationships among the major groups of sharks are still not well resolved. Many conflicting results were found, including the phylogenetic status of our study interest, *Chiloscyllium*. According to Goto [14], *C. indicum* is a sister taxon of all *Chiloscyllium* species, but this was contradicted by studies by Vélez-Zuazo and Aarnarsson [21] and Karl et al. [22], which agreed that *C. punctatum* is the sister taxon of all *Chiloscyllium* species. Thus, within this study, we aim to clarify the phylogenetic relationships of *Chiloscyllium* species through the DNA sequencing of two mitochondrial DNA loci. The cytochrome *b* region was chosen because it is the most well-studied mtDNA gene in vertebrates [23–25], including sharks, for various taxa among elasmobranchs species [8, 26, 27], at the order level [19, 21, 28], at the family level [18, 21, 29], at the genus level [30], and at the species-specific level [31, 32]. In addition, 12S rRNA has been considered to be one conserved gene among taxa, that is, helpful for phylogenetic analysis [33]. Many researchers have agreed on the important role of phylogenetic position and evolutionary distinctiveness in establishing conservation priorities [34–36]. Information about the evolutionary history of the species and the status of their close relatives can impact conservation planning, especially for those already identified as under some level of threat [21, 37].

2. Materials and Methods

2.1. Sample Collection and DNA Tissue Extraction. Tissue samples of *C. griseum* (7), *C. indicum* (19), *C. punctatum* (4), *C. hasseltii* (4), and *Stegostoma fasciatum* (1) were obtained from freshly caught animals and preserved in 95% ethanol. All of the tissue samples were collected from fishermen in Penang, Kuala Terengganu, Kuantan, Pekan, Mersing, and Pulau Tinggi, Malaysia (Figure 1). Six DNA sequences were obtained from GenBank (Table 1). Mitochondrial DNA was isolated from 25 mg of muscle tissue using a DNeasy Tissue Kit (QIAGEN Inc.), following the manufacturer’s suggested protocol.

2.2. DNA Amplification. A pair of elasmobranchs-specific set of primers, GluDG (5’-TGA CTT GAA RAA CCA YCG TTG-3’) and C61121H (5’-CTC CAG TCT TCG RCT TAC AAG-3’) [27], were used in the PCR for cytochrome *b* amplification. Thermal cycling for the cytochrome *b* locus consisted of an initial step at 94°C for 5 min; 30 cycles of 94°C denaturing for 30 sec, 47–50°C annealing for 30 sec, and 60°C elongation for 1 min; and 72°C post elongation for
7 min to complete the synthesis of elongated DNA molecules. The DNA extract (3 μl) PCR was carried out in a final volume of 20 μl reaction mixture consisting of 50–100 ng DNA template, 5 U/μl Taq DNA polymerase (PROMEGA), 5X PCR buffer, 25 mM MgCl₂, 10 mM dNTPs, and 10 pmol/μl of each primer and ddH₂O. Universal primers for the 12S rRNA gene (forward-5'-AAA CTG GGA TTA GAT ACC CTA T-3', reverse-5'-GAG GGT GAC GGG CGG TGT GT-3') as described by Kocher et al. [38] were used with suitable modifications. Thermocycler conditions were as follows: 3 min at 95°C for initial denaturation, followed by 30 cycles of amplification (1 min at 95°C, 45 sec at 56°C, and 1 min at 72°C), and a final extension for 7 min at 72°C. Each 25 μl reaction mixture contained 5 U/μl Taq DNA polymerase (Vivantis), 10X PCR buffer, 50 mM MgCl₂, 10 mM dNTPs, and ddH₂O. Amplified products were analyzed by electrophoresis in 1.5% agarose gel with ethidium bromide stain. PCR products were purified using a GF-1 PCR Clean-Up Kit (Vivantis), following the manufacturer’s protocol. All the successfully amplified PCR products were sent to an accredited laboratory, 1st Base Laboratories Sdn Bhd, for sequencing services.

2.3. Data Analysis. DNA sequence editing was performed with BioEdit version 7.0.2 and visually checked. The character-based method was used to infer phylogeny for Chiloscyllium species. Unweighted maximum parsimony (MP) analysis was used with a heuristic search to obtain the most parsimonious MP tree. This phylogenetic analysis was conducted using Phylogeny Analysis Using Parsimony (PAUP) version 4.0 [39]. Bootstrap analysis was performed using 1000 replications to assess robustness of branching patterns and analysis reliability.

Model test 3.7 [40] was used to choose the substitution model that best fits the data using the Akaike information criterion (AIC) for Bayesian analysis. Bayesian analyses were conducted using MrBayes 3.1 for both 12S rRNA and cytochrome b regions. The most appropriate model for 12S rRNA was found to be TrN+I (−lnL = 1131.1022; AIC = 2274.2043). The estimated base frequencies were as
follows: A, 0.3303; C, 0.2764; G, 0.1937; and T, 0.1997. Estimated substitution rates among these nucleotides were 1.0000 for A/C, A/T, C/G, and G/T; 2.4579 for A/G; and 5.9223 for C/T and the estimated proportion of invariable site was 0.6192 with equal rates. Bayesian inference was made by running two simultaneous metropolis-coupled MCMC chains for 380,000 generations with 0.009887 split frequency probability. Trees were sampled every 100 generations and a consensus tree was generated from 2850 trees by omitting the first 950 trees. For the cytochrome b sequences, the best model computed for Bayesian analysis was TrN+I (−lnL = 5441.2451; AIC = 10894.4902). The base frequency estimates were as follows: A, 0.3087; C, 0.2734; G, 0.1203; and T, 0.2976. Estimated substitution rates among these nucleotides were 1.0000 for A/C, A/T, C/G, and G/T; 2.6360 for A/G; and 4.7907 for C/T. Bayesian inferences were run for 550,000 generations (0.009769 split frequency probability) and a consensus tree was constructed from 4125 trees with 1375 burn-in.

12S rRNA and cytochrome b data sets were combined to construct one consensus tree. The incongruence-length difference (ILD)/partition homogeneity test was conducted to test the partition incongruence to overcome the issue of combining those two data sets [41]. These combined data sets were analyzed for MP analysis with heuristic search and 1000 replications of bootstrap analysis, using PAUP version 4.0 [39] and Bayesian inference (BI) under Akaike information criterion (AIC) framework. A nucleotide substitution model, TrN+I (−lnL = 6263.0845; AIC = 12538.1689), was selected using Model_test 3.7 [40] for BI with estimated base frequency as follows: A = 0.3104; C = 0.2740; G = 0.1389; T = 0.2768 and estimated substitution rates as follows: 1.0000 for A/C, A/T, C/G, and G/T; 2.1965 for A/G; and 4.2050 for C/T, and proportion of invariable sites (I) was 0.4677 with equal rates. A consensus tree was constructed from 1575 trees and 525 burn-in through 210,000 generations (trees were sampled every 100 generations) and 0.008094 split frequency probability.

3. Results

3.1. Data Analyses and Phylogenetic Resolution

3.1.1. Maximum Parsimony (MP). A total of 381 characters were sequenced for the 12S rRNA region; 302 were constant characters, 38 variable characters were parsimony-uninformative, and 41 characters were parsimony-informative. Maximum parsimony analysis resulted in one most parsimonious tree (Figure 2) with 109 steps, consistency index (CI) = 0.8349, homoplasy index (HI) = 0.1651, and retention index (RI) = 0.9122. Maximum parsimony analysis resulted in one most parsimonious tree (Figure 2) with 109 steps, consistency index (CI) = 0.8349, homoplasy index (HI) = 0.1651, and retention index (RI) = 0.9122 for 12S rRNA sequences. Two distinct branches divided Chiloscyllium species into two main clades: Clade A, consisting of C. indicum, C. plagiosum, and C. punctatum individuals, and Clade B, consisting of C. griseum and C. hasseltii individuals. C. indicum formed its own clade, with 92% bootstrap support value. With 97% bootstrap support, C. plagiosum became a sister taxon of C. indicum. C. punctatum individuals were grouped into their
own clade with 71% bootstrap support and became a sister taxon to *C. indicum* and *C. plagiosum*. The three individuals of *C. hasseltii* (*C. hasseltii* 1, *C. hasseltii* 3, and *C. hasseltii* 4) formed a clade with 70% bootstrap support. However, one *C. hasseltii* sample (*C. hasseltii* 2) fell into the polytomy of *C. griseum* group, forming a nonmonophyletic clade with 81% bootstrap support.

For the cytochrome *b* region, a total of 1120 characters were aligned in MP analysis; 596 characters were constant, 172 variable characters were parsimony-uninformative, and 352 characters were parsimony-informative. One most parsimonious tree was recovered, with 893 tree length, CI = 0.7436, HI = 0.2564, and RI = 0.8842 (Figure 3). As with the 12S rRNA tree topology, *Chiloscyllium* species was divided into two different clades: Clade A, consisting of *C. indicum*, *C. plagiosum*, and *C. punctatum*, and Clade B, consisting of *C. griseum* and *C. hasseltii*, with 65% and 100% bootstrap support, respectively. *C. plagiosum* and *C. indicum* formed their own clade, both with 100% bootstrap support, and became sister taxa of each other. *C. punctatum* also formed its own clade, with 100% bootstrap support. In another distinct clade (*C. griseum* and *C. hasseltii*), most *C. griseum* individuals displayed polytomies, while *C. hasseltii* created its own clade. The *C. griseum* and *C. hasseltii* clade was not monophyletic, as there were two *C. griseum* individuals in the *C. hasseltii* group. Clade B is a sister taxon of Clade A.

3.2. Bayesian Analysis. Bayesian analysis for 12S rRNA data set produced essentially the same topology as the MP analysis (Figure 2) and revealed a separation of two branches, with Bayesian posterior probabilities of 0.96 for a *C. indicum*, *C. plagiosum*, and *C. punctatum* group (Clade A) and 0.71 for a *C. griseum* and *C. hasseltii* group (Clade B). A 1.00 Bayesian posterior probability supports the clustering of the *C. punctatum* branch. *C. indicum* and *C. plagiosum* became a sister clade with 0.92 Bayesian posterior probability support. Like 12S rRNA, cytochrome *b* Bayesian tree topology also formed two distinct clades (Figure 3): Clade A, consisting of *C. indicum*, *C. plagiosum*, and *C. punctatum* individuals, and...
Clade B, consisting of *C. griseum* and *C. hasseltii* individuals, with 0.92 and 1.00 Bayesian posterior probability support, respectively. A Bayesian posterior probability of 0.92 supports the clustering of the *C. indicum* and *C. plagiosum* clade. *C. punctatum* individuals formed their own clade, with 1.00 Bayesian posterior probability.

3.3. **Consensus Analysis.** ILD test simulated nonsignificant *P* value > 0.05 (0.5200), meaning that 12S rRNA and cytochrome *b* analyses should be proceeded for phylogenetic reconstruction. The MP analysis based on equally weighted total substitutions for combined data of 12S rRNA and cytochrome *b* sequences produced a tree with 893 steps, CI = 0.7828, HI = 0.2172, and RI = 0.9038. Out of 1501 characters sequenced for both regions, 32 characters were constant, 232 variable characters were parsimony-uninformative, and 337 were parsimony-informative. The results of the MP and Bayesian analysis of all data combined are shown in Figure 4. These results were broadly similar to all analysis for the 12S rRNA and cytochrome *b* regions. *Chiloscyllium* species was divided into two major sister clades that clustered *C. griseum* and *C. hasseltii* in one group (Clade B) and *C. indicum*, *C. plagiosum*, and *C. punctatum* clade in the other (Clade A). The *C. griseum* and *C. hasseltii* nonmonophyletic clade was supported with 100% bootstrap and 1.00 Bayesian posterior probability. *C. indicum*, *C. plagiosum*, and *C. punctatum* clustered into their own clade, with 100% bootstrap support and 1.00 Bayesian posterior probability. *C. indicum* and *C. plagiosum* grouped together, with 81% bootstrap support and 1.00 Bayesian posterior probability. *C. punctatum* became sister taxon of these two species, with 93% bootstrap support and 1.00 Bayesian posterior probability.

4. **Discussion**

This molecular study has led to a better understanding of the relationships among *Chiloscyllium*. Tree topologies of the 12S rRNA region show similarities to the cytochrome *b* tree topologies for all of the analyses conducted. *C. indicum* formed its own monophyletic group and became a sister taxon of *C. plagiosum*, while *C. punctatum* group became a sister taxon of the *C. indicum* and *C. plagiosum* clade. The nonmonophyly within *C. griseum* and *C. hasseltii* was observed using the two different molecular markers for all analyses. However, it can be differentiated into two groups: (1) *C. griseum* group and (2) *C. hasseltii* group. Such grouping actually reflects the close relationships between these two species. Vélez-Zuazo and Agnarsson [21] and Naylor et al. [42] also indicated these two species as sister species.
According to Dingerkus and DeFino [5], *C. hasseltii* adults look almost identical to *C. griseum* adults; however, they can still be distinguished by morphometric measurements. For *C. griseum*, the distance between the first and second dorsal fins is usually more than 9.3% of total length (less than 9.3% in *C. hasseltii*); the height of the first dorsal fin is usually more than 6.6% of total length (less than 6.6% in *C. hasseltii*); and the height of the second dorsal fin is usually more than 5.8% of total length (less than 5.8% in *C. hasseltii*). *C. hasseltii* juveniles are separable from *C. griseum* by having black bands outlined pattern, whereas *C. griseum* does not have the bands outlined in black [5]. Therefore, our nonmonophyly *C. griseum/C. hasseltii* groups might occur due to some missing data and uneven taxon sampling which affect our topology inadequacies. Furthermore, according to Heist [43], coastal and benthic sharks like *C. griseum* and *C. hasseltii* exhibit only a little divergence in nuclear and mitochondrial gene frequencies where they are continuously distributed along continental margins.

All of the research results showed that *Chiloscyllium* species is separated into two distinctive clades: clade *C. griseum* and *C. hasseltii* (Clade B) and its sister clade, the *C. indicum, C. plagiosum*, and *C. punctatum* clade (Clade A). *C. punctatum* is a sister taxon of *C. indicum* and *C. plagiosum*. This shows that *C. indicum* is having the closest genetic relationships with *C. plagiosum* while their close sister is *C. punctatum*. *C. griseum* is the close sister to *C. hasseltii*. This finding is contrary to the study conducted by Vélez-Zuazo and Aagnarsson [21], which identified *C. indicum* and *C. plagiosum* as sisters, *C. griseum* and *C. hasseltii* as sisters, and *C. punctatum* as sister of all the other *Chiloscyllium* species. However, Vélez-Zuazo and Aagnarsson’s [21] suggested topology was questionable because the authors themselves noted that their analysis suffered from a substantial amount of missing data (85%) and their data came directly from GenBank [42] which might bear some errors and inaccuracies. Karl et al. [22] proposed that *C. indicum* is a sister of *C. plagiosum, C. hasseltii* is a sister of those two species, and *C. punctatum* is a sister of all the *Chiloscyllium* species while Goto [14] placed *C. indicum* as a sister species of all the other *Chiloscyllium* species based on their morphological synapomorphies in mode of reproduction, maximum number of offspring, and existence of vestibular horns or silky fibers on egg case. Our molecular results agreed with the hypothesis suggested by Naylor et al. [42] inferred from a Bayesian analysis based on NADH2 sequence data. *C. indicum* and *C. plagiosum* are sisters to each other and their close species is *C. punctatum*, while *C. griseum* and *C. hasseltii* formed their own respective clade. The phylogenetic placement of the specimen of *Hemiscyllium ocellatum* was found to be model dependent [42]. Dingerkus and DeFino [5] cladogram based on morphological characters was also supported by our study hypothesis. According to Dingerkus and DeFino [5], genus *Chiloscyllium* can be defined as monophyletic by possessing less than 180 vertebral, strong middorsal ridges that extend from the back of the head onto the tail, fourth and fifth gill openings that meet dorsally and ventrally, and a projection on the posterior border of the spiracular opening, pointing into the spiracle. Dingerkus and DeFino [5] indicated that *C. plagiosum* and *C. indicum* are sister species by sharing the following morphology: lateral dermal ridges on the body; specially modified dermal denticles on these dermal ridges; dermal denticles covering the rest of the body; and gill arches that are of the primitive *Chiloscyllium* type, except that *C. indicum* has lost the small basibranchial element. Dingerkus and DeFino [5] also positioned *C. punctatum* as the closest species to the *C. plagiosum* and *C. indicum* sister-species clade. It is followed by *C. griseum, C. hasseltii*, and two new species, *C. confusum* and *C. burmensis*. These remaining species form a group based on the three hypobranchial elements projecting backward to almost (or fully) touch the basibranchial plate, the basibranchial element being greatly reduced (or absent) and the adults being uniform in color dorsally, having lost any juvenile color or patterns [5]. The *C. hasseltii*, *C. griseum*, *C. confusum*, and *C. burmensis* group is defined by having the third hypobranchials articulating with the basibranchial plate, the absence of basibranchial elements, and dermal denticles as broad as they are long. This indicates that *C. griseum* and *C. hasseltii* shared some common characteristics that placed them as sister-species taxa, as shown by the present findings. *C. hasseltii* is the sister group of *C. confusum* and *C. burmensis*, based on the presence of mid-lateral keels on the dermal denticles [5].

5. Conclusion

Our study indicated that the cytochrome b and 12S rRNA genes are reliable tools to investigate the genetic relationships among five species of bamboo sharks. This study could be improved by increasing taxon sampling and adding study regions from nuclear genes to gain more informative phylogenetic estimation. These results should be considered as initial step toward understanding *Chiloscyllium* relationships as they were inferred analysis from two fast-evolving mitochondrial genes. This deposition of the *Chiloscyllium* species will pave the way for future discussion on shark evolution, conservation strategies, fisheries management, and benefit as a preliminary reference toward a better conservation assessment.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

Acknowledgments

The authors acknowledge Universiti Kebangsaan Malaysia and the Southeast Asian Fisheries Development Center (SEAFDEC), Malaysia, for providing necessary funding, facilities, and assistance. This research was supported by Grants UKM-GUP-PLW-08-11-046, BKBP-FST-K006230, and DLP-2013-006.
References

[1] J. Müller and F. G. J. Henle, Systematische Beschreibung Der Plagiostomen, Viet, Berlin, Germany, 1838.

[2] E. T. Bennett, "Class pisces," in Memoir of the Life and Public Services of Sir Thomas Stamford Raffles, S. R. Lady, Ed., pp. 686–694, John Murray Publishers, London, UK, 1830.

[3] P. Bleeker, "Bijdrage tot de kennis der Plagiostomen van den Indischen Archipel," Verhandelingen Van Het Bataviaasch Genootschap Van Kunsten En Wetenschappen, vol. 24, no. 12, pp. 1–92, 1852.

[4] Department of Fisheries Malaysia, Malaysia National Plan of Action For the Conservation and Management of Shark, Ministry of Agriculture and Agro-based Industry Malaysia, Putrajaya, Malaysia, 2006.

[5] G. Dingerkus and T. C. DeFino, "A revision of the Orectolobiform shark family Hemiscyllidae (Chondrichthyes, Selachii)," Bulletin of the American Museum of Natural History, vol. 176, no. 1, pp. 1–94, 1983.

[6] L. J. V. Compagno, FAO Species Catalogue, Vol 4. Sharks of the World, An Annotated and Illustrated Catalogue of Shark Species Known To Date, Part 1, Hexanchiformes To Lamniformes, FAO Fish Synop 125(4/1), FAO, Rome, Italy, 1984.

[7] IUCN, 2013, IUCN red list of threatened species, http://www.redlist.org/.

[8] G. J. P. Naylor, J. A. Ryburn, O. Fedrigo, and J. A. Lopez, "Phylogenetic relationships among the major lineages of modern elasmobranchs," in Reproductive Biology and Phylogeny of Chondrichthyes: Sharks, Batoids and Chimaeras, W. C. Hamlett, Ed., pp. 1–25, Science Publishers, Enfield, NH, USA, 2005.

[9] J. Müller and F. G. J. Henle, "Ueber die gattungen der plagiostomen," Archiv für Naturgeschichte, vol. 3, pp. 394–401, 1837.

[10] E. R. Waite, "Studies in Australian sharks, with a diagnosis of a new family," Records of the Australian Museum, vol. 4, no. 1, pp. 28–35, 1901.

[11] C. T. Regan, "A revision of the sharks of the family Orectolobidae," in Proceedings of the Zoological Society of London, pp. 347–364, 1908.

[12] G. P. Whitley, "Sharks of the Australasian region," Australian Journal of Zoology, vol. 14, pp. 173–187, 1967.

[13] L. J. V. Compagno, "Interrelationship of living elasmobranchs," in Interrelationships of Fishes, Greenwood et al., Ed., Academic Press, New York, NY, USA, 1973.

[14] T. Goto, "Comparative anatomy, phylogeny and cladistic classification of the Order Orectolobiformes (Chondrichthyes, Elasmobranchii)," Graduate School of Fisheries Sciences, vol. 48, no. 1, pp. 1–100, 2001.

[15] K. C. Ang, M. S. Ng, K. P. Reid et al., "Skin color variation in orang asli tribes of peninsular malaysia," PLoS ONE, vol. 7, no. 8, Article ID e42752, 2012.

[16] A. Ang, A. Srivasthan, B. M. Mdl.-Zain, M. R. B. Ismail, and R. Meier, "Low genetic variability in the recovering urban banded leaf monkey population of Singapore," Raffles Bulletin of Zoology, vol. 60, no. 2, pp. 589–594, 2012.

[17] S. P. Iglesias, G. Lecoindre, and D. Y. Sellos, "Extensive paraphilies within sharks of the order Carcharhiniformes inferred from nuclear and mitochondrial genes," Molecular Phylogenetics and Evolution, vol. 34, no. 3, pp. 569–583, 2005.

[18] J. A. López, J. A. Ryburn, O. Fedrigo, and G. J. P. Naylor, "Phylogeny of sharks of the family Triakidae (Carcharhiniformes) and its implications for the evolution of carcharhiniform placental viviparity," Molecular Phylogenetics and Evolution, vol. 40, no. 1, pp. 50–60, 2006.

[19] B. A. Human, E. P. Owen, L. J. V. Compagno, and E. H. Harley, "Testing morphologically based phylogenetic theories within the cartilaginous fishes with molecular data, with special reference to the catshark family (Chondrichthyes; Scyliorhinidae) and the interrelationships within them," Molecular Phylogenetics and Evolution, vol. 39, no. 2, pp. 384–391, 2006.

[20] M. P. Heinicke, G. J. P. Naylor, and S. B. Hedges, "Cartilaginous fishes (Chondrichthyes)," in The Timetree of Life, S. B. Hedges and S. Kumar, Eds., Oxford University Press, New York, NY, USA, 2009.

[21] X. Vélez-Zuazo and I. Agnarsson, "Shark tales: a molecular species-level phylogeny of sharks (Selachimorpha, Chondrichthyes)," Molecular Phylogenetics and Evolution, vol. 58, no. 2, pp. 207–217, 2011.

[22] S. A. Karl, A. L. F. Castro, and R. C. Garla, "Population genetics of the nurse shark (Ginglymostoma cirratum) in the western Atlantic," Marine Biology, vol. 159, no. 3, pp. 489–498, 2012.

[23] D. Zhu, S. Degnan, and C. Moritz, "Evolutionary distinctiveness and status of the endangered Lake Eacham rainbowfish (Melanotaenia eachamensis)," Conservation Biology, vol. 12, no. 1, pp. 80–93, 1998.

[24] V. F. Yun, M. C. Mahani, M. Lakim, A. Ampeng, and B. M. Md-Zain, "Phylogenetic relationships of leaf monkeys (Presbytis; Colobinae) based on cytochrome b and 12S rRNA genes," Genetics and molecular research : GMR, vol. 10, no. 1, pp. 368–381, 2011.

[25] M. K. Rosli, S. S. Zakaria, S. M. Syed-Shabthar et al., "Phylogenetic relationships of Malayan gaur with other species of the genus Bos based on cytochrome b gene DNA sequences," Genetics and Molecular Research, vol. 10, no. 1, pp. 482–493, 2011.

[26] A. P. Martin, G. J. P. Naylor, and S. R. Palumbi, "Rates of mitochondrial DNA evolution in sharks are slow compared with mammals," Nature, vol. 357, no. 6374, pp. 153–155, 1992.

[27] A. P. Martin and S. R. Palumbi, "Protein evolution in different cellular environments: cytochrome b in sharks and mammals," Molecular Biology and Evolution, vol. 10, no. 4, pp. 873–891, 1993.

[28] A. R. Hoelzel, "Shark fishing in fin soup," Conservation Genetics, vol. 2, no. 1, pp. 69–72, 2001.

[29] D. D. Lim, P. Motta, K. Mara, and A. P. Martin, "Corrigendum to “Phylogeny of hammerhead sharks (family Sphyridae) inferred from mitochondrial and nuclear genes” [Mol. Phylogen. Evol. 55 (2010) 572–579]," Molecular Phylogenetics and Evolution, vol. 58, no. 2, p. 414, 2011.

[30] B. W. Murray, J. Y. Wang, S.-C. Yang, J. D. Stevens, A. Fisk, and J. Svaravsson, "Mitochondrial cytochrome b variation in sleeper sharks (Squaliformes: Somniosidae)," Marine Biology, vol. 153, no. 6, pp. 1015–1022, 2008.

[31] D. D. Chapman, D. L. Abercrombie, C. J. Douady, E. K. Pikitch, M. J. Stanhope, and M. S. Shivji, "A streamlined, bi-organelle, multiplex PCR approach to species identification: application to global conservation and trade monitoring of the great white shark, Carcharodon carcharias," Conservation Genetics, vol. 4, no. 4, pp. 415–425, 2003.

[32] M. S. Shivji, D. D. Chapman, E. K. Pikitch, and P. W. Raymond, "Genetic profiling reveals illegal international trade in fins of the great white shark, Carcharodon carcharias," Conservation Genetics, vol. 6, no. 6, pp. 1035–1039, 2005.

[33] C. A. Matthee and S. K. Davis, "Molecular insights into the evolution of the family Bovidae: a nuclear DNA perspective;
[34] N. J. B. Isaac, S. T. Turvey, B. Collen, C. Waterman, and J. E. M. Baillie, “Mammals on the EDGE: Conservation priorities based on threat and phylogeny,” *PLoS ONE*, vol. 2, no. 3, article e296, 2007.

[35] A. Ø. Mooers, D. P. Faith, and W. P. Maddison, “Converting endangered species categories to probabilities of extinction for phylogenetic conservation prioritization,” *PLoS ONE*, vol. 3, no. 11, Article ID e3700, 2008.

[36] M. K. A. Rosli, S. M. F. Syed-Shabthar, P. Abdul-Patah et al., “A new subspecies identification and population study of the Asian small-clawed otter (*Aonyx cinereus*) in Malay Peninsula and Southern Thailand based on fecal DNA method,” *Scientific World Journal*, vol. 2014, Article ID 457350, 11 pages, 2014.

[37] S. M. N. Romaino, Z. A. Fazly-Ann, S. Loo - et al., “Species identification of Malaysian cattle using PCR-RFLP method,” *Genetics and Molecular Research*, vol. 13, no. 1, pp. 406–414, 2014.

[38] T. D. Kocher, W. K. Thomas, A. Meyer et al., “Dynamics of mitochondrial DNA evolution in animals: amplification and sequencing with conserved primers,” *Proceedings of the National Academy of Sciences of the United States of America*, vol. 86, no. 16, pp. 6196–6200, 1989.

[39] D. L. Swofford, *PAUP 4.0 Phylogenetic Analysis Using Parsimony (and Other Methods)*, Sinauer Press, Sunderland, Mass, USA, 2002.

[40] D. Posada and K. A. Crandall, “MODELTEST: testing the model of DNA substitution,” *Bioinformatics*, vol. 14, no. 9, pp. 817–818, 1998.

[41] F. K. Barker and F. M. Lutzoni, “The utility of the incongruence length difference test,” *Systematic Biology*, vol. 51, no. 4, pp. 625–637, 2002.

[42] G. J. P. Naylor, J. N. Caira, K. Jensen, K. A. M. Rosana, N. Straube, and C. Lakner, “Elasmobranch phylogeny: a mitochondrial estimate based on 595 species,” in *Biology of Sharks and Their Relatives*, J. C. Carrier, J. A. Musick, and M. R. Heithaus, Eds., pp. 37–53, CRC Press, Boca Raton, Fla, USA, 2nd edition, 2012.

[43] E. J. Heist, “Genetics of sharks, skates and rays,” in *Biology of Sharks and Their Relatives*, J. C. Carrier, J. A. Musick, and M. R. Heithaus, Eds., pp. 487–499, CRC Press, Boca Raton, Fla, USA, 2nd edition, 2012.